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NEWS 3 JUL 02 SCISEARCH enhanced with complete author names
NEWS \,4\, JUL 02 CHEMCATS accession numbers revised
NEWS 5 JUL 02 CA/CAplus enhanced with utility model patents from China
NEWS 6 JUL 16 CAplus enhanced with French and German abstracts
NEWS 7 JUL 18 CA/CAplus patent coverage enhanced
NEWS 8 JUL 26 USPATFULL/USPAT2 enhanced with IPC reclassification
NEWS 9 JUL 30 USGENE now available on STN
NEWS 10 AUG 06 CAS REGISTRY enhanced with new experimental property tags
NEWS 11 AUG 06 BEILSTEIN updated with new compounds
NEWS 12 AUG 06 FSTA enhanced with new thesaurus edition
NEWS 13 AUG 13 CA/Caplus enhanced with additional kind codes for granted
                patents
NEWS 14 AUG 20 CA/CAplus enhanced with CAS indexing in pre-1907 records
NEWS 15 AUG 27 Full-text patent databases enhanced with predefined
                patent family display formats from INPADOCDB
NEWS 16 AUG 27 USPATOLD now available on STN
NEWS 17 AUG 28 CAS REGISTRY enhanced with additional experimental
                spectral property data
NEWS 18 SEP 07 STN AnaVist, Version 2.0, now available with Derwent
                World Patents Index
NEWS 19 SEP 13 FORIS renamed to SOFIS
NEWS 20 SEP 13 INPADOCDB enhanced with monthly SDI frequency
NEWS 21 SEP 17 CA/CAplus enhanced with printed CA page images from
                1967-1998
NEWS 22 SEP 17 CAplus coverage extended to include traditional medicine
                patents
NEWS 23 SEP 24 EMBASE, EMBAL, and LEMBASE reloaded with enhancements
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NEWS EXPRESS 19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2, CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP), AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.

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NEWS IPC8 For general information regarding STN implementation of IPC 8

NEWS 24 OCT 02 CA/CAplus enhanced with pre-1907 records from Chemisches

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FILE CONTENT:1840 - 6 Oct 2007 VOL 147 ISS 16

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This file contains CAS Registry Numbers for easy and accurate substance identification.

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=> d L1 HAS NO ANSWERS L1 STR

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=> s l1

SAMPLE SEARCH INITIATED 15:31:54 FILE 'CASREACT'

SCREENING COMPLETE - 4253 REACTIONS TO VERIFY FROM 319 DOCUMENTS

100.0% DONE 4253 VERIFIED 50 HIT RXNS 4 DOCS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED VERIFICATIONS:

81160 TO 88960

PROJECTED ANSWERS:

4 TO 199

50 REACTIONS) 4 SEA SSS SAM L1 (

=> s l1 full

FULL SEARCH INITIATED 15:31:59 FILE 'CASREACT'

SCREENING COMPLETE - 102162 REACTIONS TO VERIFY FROM 6508 DOCUMENTS

100.0% DONE 102162 VERIFIED 221 HIT RXNS (3 INCOMP) 44 DOCS

SEARCH TIME: 00.00.02

44 SEA SSS FUL L1 (221 REACTIONS) L3

=> d his

L1

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44 S L1 FULL L3

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> ENTRY SESSION

114.00 114.21 FULL ESTIMATED COST

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FILE COVERS 1907 - 11 Oct 2007 VOL 147 ISS 16 FILE LAST UPDATED: 10 Oct 2007 (20071010/ED)

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=> s 13

44 L3

=> d l4 1-44 ibib abs hitrn

ANSWER 1 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

DOCUMENT NUMBER: ***147:323232***

TITLE: Novel method for the preparation of stavudine

polymorphic forms I and II

Di Lernia, Gianluca; Mancini, Alfredo; Mancini, INVENTOR(S):

Franco; Serra, Maurizio; Vergani, Domenico

Archimica Srl, Italy PATENT ASSIGNEE(S):

SOURCE: PCT Int. Appl., 16pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.					KIND DATE			APPLICATION NO.						DATE				
						-												
WO	2007	0990	82		A1 20070907			1	WO 2007-EP51812					20070226				
	W: AE, AG, A		AL,	AM,	ΑT,	ΑŪ,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,		
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	
		ΚP,	KR,	KZ,	LА,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	
		MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	
		RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	sv,	SY,	TJ,	TM,	TN,	TR,	TT,	
		TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	zw							
	RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	
		IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	
		CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG,	BW,	GH,	
		GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	ŪĠ,	ZM,	ZW,	AM,	ΑZ,	BY,	
		KG,	ΚZ,	MD,	RU,	TJ,	TM											

PRIORITY APPLN. INFO.:

IT 2006-MI360 A 20060228

CASREACT 147:323232 OTHER SOURCE(S):

A novel method for the prepn. of stavudine polymorphic forms I and II is AB described; it's prepd. starting from 5'-acetate-2',3'-dideoxy-5methyluridine. The 5'-acetate-2',3'-dideoxy-5-methyluridine, by reaction with catalytic amts. of sodium methoxide in a C1-C4 alc. solvent, gives crude stavudine form II. Crude stavudine form II, which doesn't need to be isolated or purified, can be converted into polymorphic form I by slurry at reflux in isopropanol.

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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ANSWER 2 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN
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2

ACCESSION NUMBER:

DOCUMENT NUMBER: ***144:163592***

Synthesis and antiviral activity of aryl TITLE:

phosphoramidate derivatives of .beta.-D- and

.beta.-L-C-5-Substituted 2',3'-didehydro-2',3'-dideoxy-

uridine bearing linker arms

Laduree, Daniel; Fossey, Christine; Delbederi, Zoica; AUTHOR(S):

Sugeac, Elena; Schmidt, Sylvie; Laumond, Geraldine;

Aubertin, Anne-Marie

CORPORATE SOURCE: U.F.R. des Sciences Pharmaceutiques, Centre d'Etudes

et de Recherche sur le Medicament de Normandie, Caen,

F-14032, Fr.

SOURCE: Journal of Enzyme Inhibition and Medicinal Chemistry

(2005), 20(6), 533-549

CODEN: JEIMAZ; ISSN: 1475-6366

Taylor & Francis Ltd. PUBLISHER:

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 144:163592

The authors have previously reported the synthesis and evaluation of potent anti-human immunodeficiency virus compds. based on .beta.-D-d4T

analogs bearing a tether attached at the C-5 position and their .beta.-L-counterparts. Initial study revealed a requirement for an alkyl side-chain with an optimal length of 12 carbons for a weak antiviral activity. As a continuation of that work, the authors have now prepd. the corresponding phosphoramidate derivs. as possible membrane-permeable prodrugs. Phosphorochloridate chem. gave the target phosphoramidates which were tested for anti-human immunodeficiency virus type 1 activity; unfortunately, they were devoid of anti-HIV activity.

REFERENCE COUNT: 35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 3 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

DOCUMENT NUMBER: ***145:419404***

Unusual olefin formation by PhSe-F trans-elimination TITLE: Du, Jinfa; Shi, Junxing; Chun, Byoung-Kwon; Hobbs, AUTHOR(S):

Ann; Hollecker, Laurent; Watanabe, Kyoichi A.

Pharmasset, Inc., Tucker, GA, USA CORPORATE SOURCE:

Nucleosides, Nucleotides & Nucleic Acids (2005), SOURCE:

24(9), 1289-1292

CODEN: NNNAFY; ISSN: 1525-7770

Taylor & Francis, Inc. PUBLISHER:

Journal DOCUMENT TYPE: LANGUAGE: English

OTHER SOURCE(S): CASREACT 145:419404

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A new approach to the synthesis of a 2',3'-didehydro-2',3'-AB dideoxynucleoside I was described. The reaction was carried out in excellent yield through unusual olefin formation by PhSe-F trans-elimination.

REFERENCE COUNT: 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:1094206 CAPLUS <<LOGINID::20071011>>

143:347399 DOCUMENT NUMBER:

TITLE: Method for preparing 2',3'-dideoxycytidine

INVENTOR(S): Lu, Jinkang; Wang, Xinhua

Peop. Rep. China PATENT ASSIGNEE(S):

Faming Zhuanli Shenqing Gongkai Shuomingshu, 8 pp. SOURCE:

CODEN: CNXXEV

DOCUMENT TYPE: Patent

Chinese LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1563030	Α	20050112	CN 2004-10014350	20040315
PRIORITY APPLN. INFO.:			CN 2004-10014350	20040315

OTHER SOURCE(S): CASREACT 143:347399

The title method comprises allowing cytidine to react with acetic AΒ anhydride to get 4-N-acetylcytidine, reacting 4-N-acetylcytidine with a satd. acetic acid soln. of hydrogen bromide with acetic anhydride as catalyst to get a bromide mixt.; reacting to get 4-N-acetyl-2',3'didehydro-2',3'-dideoxycytidine-5'-acetate with zinc copper couple as catalyst; hydrogenating in the presence of Pd/C to get

4-N-acety1-2',3'-dideoxyguanosine-5'-acetate; and reacting at the presence of methanol and triethylamine to get 2',3'-dideoxycytidine. The invention has the advantages of simple technique, easy operation, easily-accessible raw material, and high yield exceeding 90%.

L4 ANSWER 5 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:582511 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***143:78442***

TITLE: A process for the production of purine

didehydronucleoside compounds via elimination reaction

using perfluoro-alkane-sulfonyl fluoride

INVENTOR(S): Torii, Takayoshi; Onishi, Tomoyuki; Izawa, Kunisuke

PATENT ASSIGNEE(S): Ajinomoto Co., Inc., Japan SOURCE: Eur. Pat. Appl., 24 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	NO.		KIND DATE			APPLICATION NO.						DATE				
		· -														
EP 1550	665		A1 20050706			EP 2004-30327						20041221				
R:	AT, BE	E, CH,	DE,	DK,	ES,	FR,	GB,	GF	R, IT,	LI,	LU,	NL,	SE,	MC,	PT,	
	IE, SI	, LT,	LV,	FI,	RO,	MK,	CY,	ΑI	, TR,	BG,	CZ,	EE,	HU,	PL,	SK,	
	BA, HF	R, IS,	YU													
JP 2005	206585		Α		2005	0804		JP	2004-	3683	51		2	0041	220	
US 2005	A1		2005	0804		US	2004-	1674	1		2	0041	221			
PRIORITY APP						JP	2003-	4340	09		A 2	0031	226			
OTHER COURCE	CACI	סעמכ	TT 14	3.78	442.	ΜZ	ידעססע	143.	7844	2						

OTHER SOURCE(S): CASREACT 143:78442; MARPAT 143:78442

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The invention provides a process wherein a 3'-deoxy-3'-halopurine nucleoside compd. is treated with perfluoroalkanesulfonyl fluoride in the presence of a base to give a 2',3'-didehydro-2',3'-dideoxy-purine nucleosides. The resulting compd. may be subjected to a catalytic hydrogenolysis to obtain a 2',3'-dideoxypurine nucleoside I, wherein B is purine nucleobase; R is a protective group. Thus, 5'-O-acetyl-2',3'-dideoxyadenosine was prepd. from 5'-O-acetyl-3'-deoxy-3'.beta.-bromoadenosine via elimination reaction of 5'-O-acetyl-2',3'-didehydro-2',3'-dideoxyadenosine in good yield.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 6 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003:552720 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***140:111616***

TITLE: Synthesis of triazole nucleoside derivatives

AUTHOR(S): Li, Zicheng; Chen, Shuhua; Jiang, Ning; Cui, Gang

CORPORATE SOURCE: Faculty of Chemistry, Sichuan University, Sichuan,

Peop. Rep. China

SOURCE: Nucleosides, Nucleotides & Nucleic Acids (2003),

22(4), 419-435

CODEN: NNNAFY; ISSN: 1525-7770

PUBLISHER: Marcel Dekker, Inc.

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 140:111616

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/ Structure 14 in file .gra /

5'-O-Mesyl-2',3'-O-isopropylidene ribonucleosides were converted to their 5'-substituted nucleosides in good yields by reacted with NaN3 or KI.
2',3'-O-Isopropylidene ribonucleosides were prepd. in good yields from ribonucleosides I (R = CONH2, NO2; R1 = OH) with a reaction mixt. of acetone and tri-Et orthoformate instead of using acetone di-Et acetal. Compds. I were treated with 2-acetoxyisobutyryl halide (Cl or Br) to give 1-[2-O-acetyl-3-halo-3-deoxy-5-O-(2,5,5-trimethyl-1,3-dioxolan-4-on-2-yl)-.beta.-D-xylofuranosyl]-1,2,4-triazole-3-carboxamide in high yields using 2-acetoxyisobutyryl bromide, the mixt. of 2-acetoxyisobutyryl chloride and NaBr. Triazole nucleosides, e.g. I (R = NH2, R1 = H, NH2) were prepd. and tested as antiviral and antitumor agents. The biol. activity of title compds. was examd. in human liver cancer cells (A-549), lung cancer cells (BEL-7402), and Flu-A cells.

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 7 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003:502075 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***139:214647***

TITLE: 2'-Chloro-2',3'-dideoxy-3'-fluoro-D-ribonucleosides:

Synthesis, Stereospecificity, Some Chemical Transformations, and Conformational Analysis

AUTHOR(S): Mikhailopulo, Igor A.; Pricota, Tamara I.; Sivets,

Grigorii G.; Altona, Cornelis

CORPORATE SOURCE: Institute of Bioorganic Chemistry, National Academy of

Sciences, Minsk, 220141, Belarus

SOURCE: Journal of Organic Chemistry (2003), 68(15), 5897-5908

CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 139:214647

The synthesis of Me 5-0-benzoyl-2-chloro-2,3-dideoxy-3-fluoro-.beta.-Dribofuranoside and its use as a glycosylating agent for persilylated thymine, N6-benzoyladenine, and N4-benzoylcytosine are described. 2'-chloro-2',3'-dideoxy-3'-fluoro-D-ribonucleosides synthesized were transformed to 2',3'-dideoxy-3'-fluoro-.alpha.- and -.beta.-D-erythropentofuranoside nucleosides of thymine, adenine, and cytidine by treatment with tributyltin hydride in the presence of .alpha.,.alpha.'azobisisobutyronitrile. Treatment of 2'-chloro-2',3'-dideoxy-3'-fluoro-Dribonucleosides with 1 M MeONa/MeOH under reflux for 1-5 h afforded 2',3'-didehydro-2',3'-dideoxy-2'-chloro-D-pentofuranosyl nucleosides as the principal products (47-81%) of the reaction, along with recovered starting nucleoside. Easy HF elimination was also obsd. in the case of the 2'-azido-2',3'-dideoxy-3'-fluoro-.beta.-D-ribofuranosides of thymine and adenine. The role of conformational peculiarities of 2'-chloro-2',3'-dideoxy-3'-fluoro-D-ribonucleosides in the obsd. exclusive elimination of HF is discussed. The conformational anal. of a rather broad palette of 2,3-dideoxy-3-fluoro-2-(X-substituted)-D-ribofuranosides was performed with the aid of the PSEUROT (version 6.3) program, using (i) the recently re-parametrized Karplus-type relation (Chattopadhyaya, J. Org. Chem. 1998, 63, 4967) and (ii) empirical bond angle correction terms suggested by us. The predictive power of the Brunck and Weinhold model (J. Am. Chem. Soc. 1979, 101, 1700) of the gauche effect between atoms and groups as a conformational driving force acting upon the pentofuranose ring is explored.

REFERENCE COUNT: 73 THERE ARE 73 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 8 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003:389753 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***139:345311***

TITLE: Synthesis and Anti-HIV Activity of Some Heterodimers

[NRTI]-Glycyl-Succinyl-[Trovirdine Analogue] of Known

HIV-1 Reverse Transcriptase Inhibitors

AUTHOR(S): Sugeac, Elena; Fossey, Christine; Laduree, Daniel;

Schmidt, Sylvie; Laumond, Geraldine; Aubertin,

Anne-Marie

CORPORATE SOURCE: Centre d'Etudes et de Recherche sur le Medicament de

Normandie, Pharmaceutiques, U.F.R. des Sciences, Caen,

F-14032, Fr.

SOURCE: Journal of Enzyme Inhibition and Medicinal Chemistry

(2003), 18(2), 175-186

CODEN: JEIMAZ; ISSN: 1475-6366

PUBLISHER: Taylor & Francis Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 139:345311

Expected for their ability to inhibit HIV replication, four heterodimers with a Nucleoside Reverse Transcriptase Inhibitor (NRTI) and a Non-Nucleoside Reverse Transcriptase Inhibitor (NNRTI) bound by a linker arm were designed and synthesized. For the NRTIs, d4U, d2U, d4T and 5'-O-acetyl-5-(3-hydroxypropynyl)d2U were chosen. For the NNRTI, a Trovirdine Analog (belonging to the phenethylthiazolylthiourea class) was chosen. The conjugation of the two different inhibitors (NRTI and NNRTI) was performed using the succinyl-glycine moiety as a spontaneously cleavable linker.

REFERENCE COUNT: 34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 9 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003:123466 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***138:137526***

TITLE: Preparation of 2',3'-didehydro-3'-deoxythymidine

INVENTOR(S): Yao, Qizheng; Lu, Gang

PATENT ASSIGNEE(S): China Pharmacy Univ., Peop. Rep. China

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 12 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1324800	A	20011205	CN 2001-113661	20010601
PRIORITY APPLN. INFO.:			CN 2001-113661	20010601

OTHER SOURCE(S): CASREACT 138:137526; MARPAT 138:137526

AB Stavudine (2',3'-didehydro-3'-deoxythymidine) as antiviral agent is synthesized by dehydrating 5-methyluridine with di-C1- 4 alkyl or di-Ph carbonate (at a molar ratio of 1.1-1.5) in anhyd. polar solvent (such as DMF or DMA) in the presence of base (such as NaHCO3 or KHCO3) at 90-150.degree. to obtain 2,2'-anhydro-5-methyluridine; acylating and halogenating with C1- 6 acyl halide (or org. anhydride-org. acid-NaBr and/or H2SO4, or org. anhydride-pyridinium bromide perbromide) in polar solvent (such as acetonitrile, DMF, or Et acetate) at 50-75.degree. to

obtain 2'-halo-3',5'-O-diacylthymidine (halo = Cl, Br, or I); reductive eliminating with metal reductant (such as Zn or Zn-Cu) in methanol or ethanol and/or Et acetate in the presence of acid (such as acetic acid or propanoic acid) at 5-35.degree.; and hydrolyzing with base (such as Na methoxide or NH3) in methanol or ethanol and/or dioxane.

ANSWER 10 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

DOCUMENT NUMBER: ***137:217180***

Method for the synthesis of 2',3'-dideoxy-2',3'-TITLE:

didehydronucleosides

Jin, Fuqiang; Confalone, Pasquale N. INVENTOR(S):

Pharmasset Ltd., USA PATENT ASSIGNEE(S): PCT Int. Appl., 180 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent

English LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA'	TENT	NO.			KIND DATE			APPLICATION NO.						DATE				
	2002																	
WO	2002	0705	33		A3		2002	1219										
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB	, BG,	BR,	BY,	ΒZ,	CA	, CH,	CN,	
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		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ	, GW,	ML,	MR,	NE,	SN	, TD,	TG	
CA	2439				A1			CA 2002-2439836										
AU	2002	2556	54		A1	A1 20020919 AU 2002-255654							54	20020301				
	2002						2002	1226		US	2002-	8711	2			20020	301	
US	6927	291			B2		2005											
EP	1363	927			A2		2003	1126		ΕP	2002-	7250	67			20020	301	
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE	, MC,	PT,	
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL	, TR							
CN	1505	635			Α		2004	0616			2002-					20020	301	
	2002					2004	0629							:	20020	301		
JP	2004	5275	04		T		2004	0909	JP 2002-569852									
NZ	5409	56			Α		2007	0126	NZ 2002-540956						20020301			
ZA	2003	0067	39		A		2004	0901	ZA 2003-6739						20030828			
MX	2003	PA07	853		A		2004	0524								20030	901	
IN	2003	DN01	529		A		2006	1222		IN	2003-	DN15	29			20030	924	
US	2005	2509	46		A1		2005	1110		US	2005-	1828	62			20050	718	
IN	2005	DN04	086		Α		2007	0824		IN	2005-	DN40	86			20050	912	
IN	2005	DN04	087		Α		2007	0824		IN	2005-	DN40	87			20050	912	
IN	2005	DN04	880		Α		2007	0824		IN	2005-	DN40	88			20050	912	
IN	2005	DN04	090		Α		2007	0824		IN	2005-	DN40	90			20050	912	
PRIORIT	Y APP	LN.	INFO	.:						US	2001-	2724	34P		P	20010	301	
										US	2001-	2724	41P		P	20010	301	
										US	2002-	8711	2		A3	20020	301	
										WO	2002-	US64	60		W	20020	301	
										IN	2003-	DN15	29		A 3	20030	924	
OTHER S	THER SOURCE(S):					CASREACT 137:217180; MARPAT 137:217180												

The present invention is an efficient synthetic route to 2',3'-dideoxy-2',3'-didehydro-nucleosides from available precursors with the option of introducing functionality as needed, such as, the 2',3'-dideoxy- and 2'- or 3'-deoxyribo-nucleoside analogs as well as

addn1. derivs. obtained by subsequent functional group manipulations. Briefly, the present invention discloses a method for the prepn. of .beta.-D- and .beta.-L-2',3'-dideoxy-2',3'-didehydro-nucleosides starting from appropriately substituted ribonucleosides in two, optionally three steps: step (1) a halo-acylation, such as halo-acetylation, and in particular, bromo-acetylation; step (2) a reductive elimination; and optionally, step (3) a deprotection. The halo-acylation of step (1) can form the 2'-acyl-3'-halo-nucleoside, the 3'-acyl-2'-halo-nucleoside, or a mixt. thereof.

L4 ANSWER 11 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2002:325330 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***138:73449***

TITLE: Synthesis of Stavudine

AUTHOR(S): Jin, Li-ren; Jiang, Hong-ping; Hou, Peng-yi

CORPORATE SOURCE: Dept. of Chem., Xiamen Univ., Xiamen, 361005, Peop.

Rep. China

SOURCE: Xiamen Daxue Xuebao, Ziran Kexueban (2002), 41(2),

207-210

CODEN: HMHHAF; ISSN: 0438-0479

PUBLISHER: Xiamen Daxue

DOCUMENT TYPE: Journal LANGUAGE: Chinese

OTHER SOURCE(S): CASREACT 138:73449

Title compd., synthesized from ribofuranose tetraacetate and thymine via the intermediate 5-methyluridine, was described. Ribofuranose tetraacetate was condensed with bis(trimethylsilyl)thymine followed by deacylation to give 5-methyluridine, then converted to 2',3'-olefinic nucleoside by reductive elimination of 2'-bromo-3'-mesylic ester. The final product was obtained from deprotection of the hydroxy group with total yield 46.4%.

L4 ANSWER 12 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:762991 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***135:304107***

TITLE: Process for preparing 5'-acetylstavudine by the

reductive elimination reaction of 2'-deoxy-2'-bromo-

3',5'-diacetyl-5-methyluridine

INVENTOR(S): Bertolini, Giorgio; Frigerio; Marco; Velati, Maurizio;

Petrucciani, Luigi

PATENT ASSIGNEE(S): Clariant Life Science Molecules, Italy

SOURCE: PCT Int. Appl., 12 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE		
WO 2001077103	A1 20011018	WO 2001-IB588	. 20010410		
W: BR, HU, JP,	KR, MX				
RW: AT, BE, CH,	CY, DE, DK, ES,	FI, FR, GB, GR, IE, IT,	LU, MC, NL,		
PT, SE, TR					
IT 2000MI0810	A1 20011012	IT 2000-MI810	20000412		
IT 1318463	B1 20030825				
EP 1274704	A1 20030115	EP 2001-917385	20010410		
EP 1274704	B1 20040114				
R: AT, BE, CH,	DE, DK, ES, FR,	GB, GR, IT, LI, LU, NL,	SE, MC, PT,		
IE, FI, CY,	TR				
BR 2001009965	A 20030527	BR 2001-9965	20010410		
HU 200300259	A2 20030628	HU 2003-259	20010410		

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T
                                                          JP 2001-575576
      JP 2003530394
                                          20031014
                                                                                         20010410
                                T 20040115 AT 2001-917385
T3 20040816 ES 2001-1917385
A1 20020226 US 2001-834028
A 20030310 MX 2002-PA10072
      AT 257835
                                                                                         20010410
      ES 2213109
                                                                                         20010410
      US 2001039342
                                                                                       20010412
                                                                                       20021011
      MX 2002PA10072
                                                          IT 2000-MI810 A 20000412 WO 2001-IB588 W 20010410
PRIORITY APPLN. INFO.:
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OTHER SOURCE(S): CASREACT 135:304107

5'-Acetylstavudine is prepd. in high yield and selectivity by the reaction of 2'-deoxy-2'-bromo-3',5'-diacetyl-5-methyluridine in the presence of zinc as reducing agent combined with an activating agent which is an ammonium salt (e.g., triethylammonium hydrochloride) and/or a phosphonium salt in a dipolar aprotic solvent (e.g., DMF).

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 13 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:735911 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***136:37877***

TITLE: Synthesis of 9-(2,3-Dideoxy-2-fluoro-.beta.-D-threo-

pentofuranosyl) adenine (FddA) via a Purine

3'-Deoxynucleoside

AUTHOR(S): Takamatsu, Satoshi; Maruyama, Tokumi; Katayama,

Satoshi; Hirose, Naoko; Naito, Masaki; Izawa, Kunisuke

CORPORATE SOURCE: AminoScience Laboratories, Ajinomoto Co., Inc.,

Suzuki-cho, Kawasaki-ku, Kawasaki, Kanagawa, 210-8681,

Japan

SOURCE: Journal of Organic Chemistry (2001), 66(22), 7469-7477

CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 136:37877

A synthesis of 9-(2,3-dideoxy-2-fluoro-.beta.-D-threo-pentofuranosyl)adenine (FddA) via a 6-chloro-9-(3-deoxy-.beta.-D-erythro-pentofuranosyl)-9H-purine, which was readily obtained from inosine, is described. Fluorination at the C2'-.beta. position of the purine 3'-deoxynucleoside with diethylaminosulfur trifluoride was improved by the introduction of a 6-chloro group and proceeded in moderate yield. Purine 3'-deoxynucleoside derivs. were also subjected to nucleophilic reactions with triethylamine trihydrofluoride and gave the desired fluorinated nucleoside in good yield. The safety and yield of the fluorination process were greatly improved by the use of triethylamine trihydrofluoride. The influence of the sugar ring conformation and 6-chloro group on the rate of the nucleophilic reaction against elimination are also discussed.

REFERENCE COUNT: 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 14 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:710913 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***136:86017***

TITLE: Synthesis of 5-alkenylated D4T analogs via the

Pd-catalyzed cross-coupling reaction

AUTHOR(S): Ciurea, A.; Fossey, C.; Benzaria, S.; Gavriliu, D.;

Delbederi, Z.; Lelong, B.; Laduree, D.; Aubertin, A.

M.; Kirn, A.

CORPORATE SOURCE: Centre d'Etudes et de Recherche sur le Medicament de

Normandie U.F.R. des Sciences Pharmaceutiques, Caen,

14032, Fr.

SOURCE: Nucleosides, Nucleotides & Nucleic Acids (2001),

20(9), 1655-1670

CODEN: NNNAFY; ISSN: 1525-7770

PUBLISHER: Marcel Dekker, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 136:86017

The target compds. 5-[N-(6-amino-hexyl)-acrylamide]-2',3'-didehydro-2',3'-dideoxy'uridine (I) and 5-{N-[5-(methoxycarbonyl)-pentyl]-acrylamide}-2',3'-didehydro-2',3'-dideoxy-uridine (II) were prepd. by the palladium acetate-triphenylphosphine-catalyzed reaction of the 5'-0-acetyl-5-iodo-d4T analog. These compds. I and II can be used to prep. nucleotide probes carrying fluorescent labels and were nevertheless screened for their anti-HIV activity. The biol. data demonstrated that none of them were active against HIV-1.

REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 15 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:675236 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***136:118684***

TITLE: Synthesis of haptens for the development of

immunoassays for the monitoring of intracellular

anti-HIV nucleosides and nucleotides

AUTHOR(S): Brossette, T.; Klein, E.; Creminon, C.; Grassi, J.;

Mioskowski, C.; Lebeau, L.

CORPORATE SOURCE: Faculte de Pharmacie, Laboratoire de Synthese

Bioorganique, Universite Louis Pasteur de Strasbourg,

CNRS, Illkirch, 67 401, Fr.

SOURCE: Tetrahedron (2001), 57(38), 8129-8143

CODEN: TETRAB; ISSN: 0040-4020

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 136:118684

GI

/ Structure 15 in file .gra /

AB A series of nine modified dideoxynucleosides, e.g. I (R1 = H), and dideoxynucleotides, e.g. I (R1 = PO3H2) has been synthesized for prepg. antigenic conjugates with keyhole lympet haemocyanin in order to produce specific antibodies, and develop immunoassays. Derivs. of ddI, ddA, d4T, 3TC, and the corresponding 5'-O-monophosphates were designed incorporating an amino spacer at the base for conjugation with the proteinic antigenic carrier.

REFERENCE COUNT: 52 THERE ARE 52 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 16 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:791143 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***134:71824***

TITLE: Synthesis and antiviral activity of C-5 substituted

.beta.-D- and .beta.-L-D4t analogues

AUTHOR(S): Delbederi, Z.; Fossey, C.; Fontaine, G.; Benzaria, S.;

Gavriliu, D.; Ciurea, A.; Lelong, B.; Laduree, D.;

Aubertin, A. M.; Kirn, A.

CORPORATE SOURCE: Centre d'Etudes et de Recherche sur le Medicament de

Normandie U.F.R. des Sciences Pharmaceutiques, Caen,

14032, Fr.

SOURCE: Nucleosides, Nucleotides & Nucleic Acids (2000),

19(9), 1441-1461

CODEN: NNNAFY; ISSN: 1525-7770

PUBLISHER: Marcel Dekker, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 134:71824

AB A series of .beta.-D-2',3'-didehydro-2',3'-dideoxy-nucleosides bearing a tether attached at the C-5 position and their .beta.-L-counterparts was synthesized. Their inhibitory activities against human immunodeficiency virus (HIV) were investigated and compared to establish relationship(s) between compd. structure and their antiviral activity. No significant activity was obsd. for most .beta.-D- and .beta.-L-modified nucleosides, but 5-[N-(12-aminododecyl)carbamoylmethyl]-1-(2',3'-dideoxy-.beta.-D-glycero-pent-2-enofuranosyl)uracil and its .beta.-L-counterpart exhibited a weak activity against HIV-1.

REFERENCE COUNT:

THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 17 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:429517 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***133:193397***

TITLE: Synthesis and antiviral activity of C-5 substituted

analogs of d4T bearing methylamino- or

methyldiamino-linker arms

AUTHOR(S): Gavriliu, D.; Fossey, C.; Fontaine, G.; Benzaria, S.;

Ciurea, A.; Delbederi, Z.; Lelong, B.; Laduree, D.;

Aubertin, A. M.; Kirn, A.

CORPORATE SOURCE: Centre d'Etudes et de Recherche sur le Medicament de

Normandie U.F.R. des Sciences Pharmaceutiques, Caen,

14032, Fr.

SOURCE: Nucleosides, Nucleotides & Nucleic Acids (2000), 19(5

& 6), 1017-1031

CODEN: NNNAFY; ISSN: 1525-7770

PUBLISHER: Marcel Dekker, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 133:193397

GI

/ Structure 16 in file .gra /

AB A general strategy is reported for the prepn. of C-5-methylamino- or methyldiamino-d4T analogs of "different sizes". Reactions of the 2',3'-didehydro-2',3'-dideoxy-C-5 hydroxymethyl precursor I(R = OH) with either polymethylene diamines NH2(CH2)nNH2COCF3 (n = 6, 8, 10 and 12) or propargylamine proceed regioselectively via substitution reactions at the C-5 position of uracil. The compds. were evaluated for antiviral activity and cytotoxicity. No significant activity was obsd. for compds. I (R = NH(CH2)nNH2COCF3; n = 9, 11, 13), but I (R = NH(CH2)nNH2COCF3; n = 10, 12) exhibited a weak activity against HIV-1.

REFERENCE COUNT: 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 18 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1999:275269 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***130:325330***

TITLE: Synthesis of L-analogs of 1-(2',3'-dideoxy-.beta.-D-

glyceropent-2-enofuranosyl)thymine

AUTHOR(S): Camara, F.; Ciurea, A.; Delbederi, Z.; Fossey, C.;

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Fontaine, G.; Gavriliu, D.; Jouenne, J.; Laduree, D.;
                         Aubertin, A. M.; Kirn, A.
CORPORATE SOURCE:
                         Centre d'Etudes et de Recherche sur le Medicament de
                         Normandie, Laboratoire de Pharmacochimie, UFR des
                         Sciences Pharmaceutiques, Caen, 14032, Fr.
                         Pharmacy and Pharmacology Communications (1999), 5(3),
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SOURCE:

225-231

CODEN: PPCOFN; ISSN: 1460-8081

Royal Pharmaceutical Society of Great Britain PUBLISHER:

DOCUMENT TYPE: Journal English LANGUAGE:

CASREACT 130:325330 OTHER SOURCE(S):

.beta.-L(-)-2',3'-Dideoxy-3'-thiacytidine (3TC), its 5-fluoro deriv. ((-)-FTC), 2',3'-dideoxycytidine (.beta.-L-ddC), and its 5-fluoro deriv. (.beta.-L-FddC) have been reported to have anti-HIV and anti-HBV activity. It was of particular interest therefore to develop a series of .beta.-L-d4T analogs bearing several kinds of amino-linker arms at the C-5 position of the pyrimidine moiety in an attempt to find more potent and less toxic anti-HIV agents. In addn., modification of nucleosides with various functional mols. has been attracting wide interest in biol. studies since the primary amino groups could be useful for the attachment of either fluorescent dyes or a non-nucleosidic reverse transcriptase inhibitor. These modified nucleosides were evaluated for antiviral activity against HIV-1LAI in CEM-SS cells and HIV-1IIIB in MT4 cells. Unfortunately, none of the compds. exhibited significant anti-HIV activity at the doses tested.

REFERENCE COUNT: 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 19 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER:

128:180621 DOCUMENT NUMBER:

Synthesis of analogs of 1-(2',3'-dideoxy-.beta.-D-TITLE:

glycero-pent-2-enofuranosyl)thymine

Fontaine, G.; Lequenne, C.; Fossey, C.; AUTHOR(S):

Renoud-Grappin, M.; Laduree, D.

Laboratoire de Pharmacochimie, Centre d'Etudes et de CORPORATE SOURCE:

Recherche sur le Medicament de Normandie, UFR des

Sciences Pharmaceutiques, Caen, 14032, Fr.

Pharmacy and Pharmacology Communications (1998), 4(1), SOURCE:

CODEN: PPCOFN; ISSN: 1460-8081

Royal Pharmaceutical Society of Great Britain PUBLISHER:

Journal DOCUMENT TYPE: English LANGUAGE:

OTHER SOURCE(S): CASREACT 128:180621

This work describes the synthesis of three new anti-HIV analogs of 1-(2',3'-dideoxy-.beta.-D-glycero-pent-2-enofuranosyl)thymine by two different routes. The compds. were functionalized on the 5-position by a carboalkoxymethyl group. The C-5 substituent on the pyrimidine ring was a

precursor for subsequent linkage with a non-nucleosidic moiety.

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 5 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 20 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

128:167643 DOCUMENT NUMBER:

A new thymine free synthesis of the anti-AIDS drug d4T TITLE:

via regio/stereo controlled .beta.-elimination of

bromoacetates

Chen, Bang-Chi; Quinlan, Sandra L.; Reid, J. Gregory; AUTHOR(S):

Spector, Richard H.

Process Exploration Labs I, Technical Operations, CORPORATE SOURCE:

Bristol-Myers Squibb Company, Syracuse, NY, 13221, USA

Tetrahedron Letters (1998), 39(8), 729-732 SOURCE:

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

CASREACT 128:167643 OTHER SOURCE(S):

The anti-AIDS drug d4T was prepd. without contamination of the nucleoside AB bond cleaved byproduct thymine from the readily available ribonucleoside 5-methyluridine. This was accomplished by using a new strategy which

involved a regio/stereo controlled .beta.-elimination of

trans-bromoacetates.

THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS 17 REFERENCE COUNT:

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 21 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN L4

ACCESSION NUMBER:

127:50881 DOCUMENT NUMBER:

Highly functionalized cyclopentanes by radical TITLE:

cyclization of unsaturated bromolactones. I.

Preparation of 5-deoxycarbahexofuranoses

Horneman, Anne Marie; Lundt, Inge AUTHOR(S):

Department of Organic Chemistry, Technical University CORPORATE SOURCE:

of Denmark, Lyngby, DK-2800, Den.

Tetrahedron (1997), 53(20), 6879-6892 SOURCE:

CODEN: TETRAB; ISSN: 0040-4020

Elsevier PUBLISHER: DOCUMENT TYPE: Journal English LANGUAGE:

CASREACT 127:50881 OTHER SOURCE(S):

Three carbasugars: 5-Deoxycarba-.alpha.-L-xylo-hexofuranose,

5-deoxycarba-.alpha.-L-lyxo-hexofuranose and 5-deoxycarba-.beta.-D-lyxo-

hexofuranose have been prepd. starting from readily available 2,7-dibromo-2,7-dideoxy-D-glycero-D-ido-heptono-1,4-lactone and 2,7-dibromo-2,7-dideoxy-D-glycero-L-gluco-heptono-1,4-lactone.

2,3-Unsatd. 7-bromo-7-deoxy-heptono-1,4-lactones were prepd. by reductive elimination of the starting compds. The key step was a highly regio- and

stereoselective 5-exo-trig radical cyclization of the unsatd.

bromolactones to give bicyclic cyclopentane derivs. The lactone moiety of these compds. were reduced using H3B - S(CH3)2 to give the above-mentioned carbahexofuranoses.

29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT:

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 22 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:109056 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***124:290128***

Synthesis of 1-(2,3-dideoxy-.beta.-D-glycero-pent-2-TITLE:

enofuranosyl) thymine (d4T; stavudine) from

5-methyluridine

Shiragami, Hiroshi; Ineyama, Takashi; Uchida, Yumiko; AUTHOR(S):

Izawa, Kinisuke

Central Res. Labs., Ajinomoto Co., Inc., Kawasaki, CORPORATE SOURCE:

210, Japan

SOURCE: Nucleosides & Nucleotides (1996), 15(1-3), 47-58

CODEN: NUNUD5; ISSN: 0732-8311

PUBLISHER: Dekker DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 124:290128

GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

A practical synthetic method of d4T (I) from 5-methyluridine (II) was AB developed. The Marumoto-Mansuri method was modified using 2',3'-O-methoxyethylidene-5-methyluridine (III) as an intermediate to afford 1-(3,5-di-O-acetyl-2-bromo-2-deoxy-.beta.-D-ribofuranosyl)thymine (IV) in high yield with less formation of byproducts. The reaction mechanism was also discussed.

ANSWER 23 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN L4

ACCESSION NUMBER: 1996:53061 CAPLUS <<LOGINID::20071011>>

124:202920 DOCUMENT NUMBER:

Allylic Substitution of 3',4'-Unsaturated Nucleosides: TITLE:

Organosilicon-Based Stereoselective Access to

4'-C-Branched 2',3'-Didehydro-2',3'-

dideoxyribonucleosides

Haraguchi, Kazuhiro; Tanaka, Hiromichi; Itoh, AUTHOR(S):

Yoshiharu; Yamaguchi, Kentaro; Miyasaka, Tadashi

School of Pharmaceutical Sciences, Showa University, CORPORATE SOURCE:

Tokyo, 142, Japan

SOURCE: Journal of Organic Chemistry (1996), 61(3), 851-8

CODEN: JOCEAH; ISSN: 0022-3263

American Chemical Society PUBLISHER:

DOCUMENT TYPE: Journal English LANGUAGE:

CASREACT 124:202920 OTHER SOURCE(S):

GI

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/ Structure 17 in file .gra /
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Reactions of organosilicon reagents (such as allyltrimethylsilane, silyl AΒ enol ethers, cyanotrimethylsilane) with 3',4'-unsatd. nucleosides, e.g. I (R = Ac, Bz, TBDPS), were investigated in the presence of a Lewis acid in CH2Cl2. In the cases of uracil and N4-acetylcytosine derivs., SnCl4 appeared to be suitable, whereas the use of EtAlCl2 was necessary for the hypoxanthine derivs. The main pathway of these reactions was found to be .alpha.-face-selective SN2' allylic substitution, irresp. of the configuration of 2'-O-acyl leaving group. As a result, a new stereoselective operation for C-C bonds formation leading to 4'-carbon-substituted 2',3'-didehydro-2',3'-dideoxyribonucleosides, e.g. II (R = Ac, Bz, TBDPS), has been disclosed for the first time. Stereochem. of these 4'-C-branched products can be assigned on the basis of 1H NMR spectroscopy in terms of the anisotropic shift of H-5 of the pyrimidine base (or H-8 of the hypoxanthine), which is caused by the 5'-O-(tert-butyldiphenylsilyl) protecting group.

ANSWER 24 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

123:144491 DOCUMENT NUMBER:

Synthesis of 5'-halo-2',3'-lyxo-epoxy and TITLE:

2',3'-unsaturated thieno[3,2-d]pyrimidine nucleosides

Rossey, Christine; Laduree, Daniel; Robba, Max AUTHOR(S):

U.F.R. des Sciences Pharmaceutiques, Centre d'Etudes CORPORATE SOURCE:

et de Recherche sur le Medicament de Normandie, Caen,

14032, Fr.

SOURCE: Journal of Heterocyclic Chemistry (1995), 32(2),

627-35

CODEN: JHTCAD; ISSN: 0022-152X

PUBLISHER: HeteroCorporation

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 123:144491

GI

/ Structure 18 in file .gra /

AB A series of thieno[3,2-d]pyrimidine-2,4-dione nucleosides, e.g. I and II, modified in the carbohydrate moiety has been synthesized. However, none of the compds. in this series exhibited significant antiviral activity against HIV at the doses tested.

L4 ANSWER 25 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1995:83000 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***122:10448***

TITLE: Nucleosides. III. Investigation of the electrochemical

synthesis of N4,05'-diacetyl-2',3'-dideoxy-2',3'-

didehydrocytidine

AUTHOR(S): Johansen, Oddvar; Marcuccio, Sebastian M.; Mau, Albert

W.-H.

CORPORATE SOURCE: Div. Chemicals Polymers, CSIRO, Clayton, 3169,

Australia

SOURCE: Australian Journal of Chemistry (1994), 47(10),

1843-50

CODEN: AJCHAS; ISSN: 0004-9425

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 122:10448

GI

/ Structure 19 in file .gra /

AB The influence of the reaction conditions on the yield of dideoxydidehydrocytidine I, by electrochem. synthesis from bromodeoxycytidine II, has been studied in order to evaluate the potential of this reaction for synthesis on a larger scale. We have characterized the half-wave potentials of the precursor II and the product I by polarog. under various conditions, and found that redn. in the base moiety can easily take place giving byproducts. Furthermore, this redn. consumes protons leading to rapid solvolysis in protic solvents. We have demonstrated for the first time that I can be formed near quant. in both protic and aprotic solvents. The success of the synthesis of I as well as of other 2',3'-dideoxy-2',3'-didehydro nucleosides also depends to a large extent on how uniform the c.d. across the working electrode surface can be maintained during electrolysis.

L4 ANSWER 26 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1995:64940 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***123:9835***

TITLE: Small-Molecule Immunostimulants. Synthesis and Activity of 7,8-Disubstituted Guanosines and

Structurally Related Compounds

AUTHOR(S): Reitz, Allen B.; Goodman, Michael G.; Pope, Barbara

L.; Argentieri, Dennis C.; Bell, Stanley C.; Burr, Levelle E.; Chourmouzis, Erika; Come, Jon; Goodman,

Jacquelyn H.; et al.

CORPORATE SOURCE: Medicinal Chemistry Department, R. W. Johnson

Pharmaceutical Research Institute, Spring House, PA,

19477, USA

SOURCE: Journal of Medicinal Chemistry (1994), 37(21), 3561-78

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 123:9835

A series of 7,8-disubstituted guanosine derivs. was designed and prepd. as AB potential B-cell-selective activators of the humoral immune response. These compds. were evaluated for their ability to act as B-cell mitogens and to augment the antibody response of B cells to sheep red blood cell (SRBC) challenge (adjuvanticity). In addn., they were tested for their ability to stimulate the natural killer (NK) cell response in murine in vitro cell assays. Certain of the compds. demonstrated in vivo activity when administered either i.v., s.c., or orally. Compds. bearing hydroxyalkyl, aminoalkyl, or substituted aminoalkyl substituents on this 7-position were weakly active. Oxo, thioxo, and seleno groups on C-8 of the quanosine ring all imparted strong activity, whereas other larger substituents did not (e.g., N:CN). A total of 80 compds. were prepd. and evaluated for their immunostimulating activity. Within this group, compds. could be divided into those that were active in all three assays, those that displayed some measure of selectivity for the adjuvanticity assay, and those that preferentially activated NK responses. Because of its overall biol. profile and ease of synthesis,7-allyl-8-oxoguanosine (loxoribine, RWJ-21757) was chosen for further development. It is among the most potent compds. evaluated in the three biol. assays.

L4 ANSWER 27 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1993:496031 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***119:96031***

TITLE: An efficient and general synthesis of 5'-esters of 2',3'-didehydro-2',3'-dideoxynucleosides: a facile

opening of 2',3'-orthoacetates of ribonucleosides

followed by reductive elimination of the halo acetates Talekar, Ratnakar R.; Coe, Paul L.; Walker, Richard T.

CORPORATE SOURCE: Sch. Chem., Univ. Birmingham, Birmingham, B15 2TT, UK

SOURCE: Synthesis (1993), (3), 303-6

CODEN: SYNTBF; ISSN: 0039-7881

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 119:96031

GI

AUTHOR(S):

/ Structure 20 in file .gra /

AB A three-step reaction sequence starting from a ribonucleoside I [B = N4-acetylcytosinyl (C), uracilyl (U), 6-oxopurinyl (I), adeninyl (A)] to give the corresponding 5'-O-acyl-2',3'-didehydro-2',3'-dideoxynucleoside II is described. The key intermediate is the bromoacetate III (B = C, I, A, R1 = Br, R2 = H, R3 = OAc; B = U, R1 = H, R2 = OAc, R3 = Br), made by reaction of the 2',3'-methoxyethylidene nucleoside IV with acetyl bromide. Reductive elimination of the bromoacetate using a zinc-copper couple

furnishes the desired compds. in good overall yield.

ANSWER 28 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1993:409106 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***119:9106***

TITLE: Nucleoside derivatives and their production

INVENTOR(S): Shiragami, Hiroshi; Uchida, Yumiko; Izawa, Kunisuke

PATENT ASSIGNEE(S): Ajinomoto Co., Inc., Japan SOURCE: Eur. Pat. Appl., 16 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE		
EP 519464	A1 -	19921223	EP 1992-110299		19920617		
EP 519464	B1	19951220					
R: BE, DE,	FR, GB, IT						
JP 05001092	A	19930108	JP 1991-245290		19910619		
JP 3042073	B2	20000515					
US 5625057	A	19970429	US 1994-295478		19940825		
PRIORITY APPLN. INFO.	:		JP 1991-245290	Α	19910619		
			US 1992-899906	B1	19920617		

OTHER SOURCE(S): CASREACT 119:9106; MARPAT 119:9106

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/ Structure 21 in file .gra /
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AB Nucleosides I [R = H, alkyl, acyl, silyl protective group; R1, R2 = alkyl; R3 = (un)protected OH, NH2; R4 = H, alkyl, halogen] were prepd. as intermediates for the unsatd. nucleosides II. Thus, 5-methyluridine was treated with MeC(OMe)3 to give 95% I [R = H, R1, R2, R4 = Me, R3 = OH (III)] as a mixt. of diastereomers. Treatment of III with 1 equiv. AcBr and 3 equiv. HBr in AcOH gave 91.0% bromouridine IV. Treatment of IV with Zn-MeCN followed by aq. Na EDTA gave 80.5% II (R = H, R3 = OH, R4 = Me).

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L4 ANSWER 29 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN
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ACCESSION NUMBER: 1992:470217 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***117:70217***

TITLE: A new synthesis of 2',3'-dideoxyinosine AUTHOR(S): Bhat, V.; Stocker, E.; Ugarkar, B. G.

CORPORATE SOURCE: Pharm-Eco Lab., Inc., Simi Valley, CA, 93065, USA SOURCE: Synthetic Communications (1992), 22(10), 1481-6

CODEN: SYNCAV; ISSN: 0039-7911

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 117:70217

AB The title compd. is prepd. in excellent yield in 4 steps starting from inosine.

L4 ANSWER 30 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1992:470212 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***117:70212***

TITLE: A convenient method for the synthesis of

2',3'-didehydro-2',3'-dideoxy nucleosides

AUTHOR(S): Dorland, Erwin; Serafinowski, Pawel

CORPORATE SOURCE: Drug Dev. Sect., Inst. Cancer Res., Sutton/Surrey, SM2

5NG, UK

SOURCE: Synthesis (1992), (5), 477-81

CODEN: SYNTBF; ISSN: 0039-7881

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 117:70212

GI

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/ Structure 22 in file .gra /
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AB 2',3'-Didehydro-2',3'-dideoxy nucleosides I (B = R, R1, adenine) were prepd. via a free radical .beta.-elimination of bromo and phenoxy(thiocarbonyl) leaving groups from appropriate phenoxy(thiocarbonyl)bromo derivs. II, adenosine, inosine, and tubercidin with Bu3SnH and subsequent deprotection of the resulting 5'-O-(2-acetoxyisobutyryl)-2',3'-didehydro-2',3'-dideoxy-nucleosides.

L4 ANSWER 31 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1991:583735 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***115:183735***

TITLE: Samarium(II) iodide-promoted deacetoxylation of

O-acetyl sugar lactones. An easy access to deoxy sugar

lactones

AUTHOR(S): Inanaga, Junji; Katsuki, Junko; Yamaguchi, Masaru

CORPORATE SOURCE: Inst. Mol. Sci., Okazaki, 444, Japan SOURCE: Chemistry Letters (1991), (6), 1025-6

CODEN: CMLTAG; ISSN: 0366-7022

DOCUMENT TYPE:

Journal English

LANGUAGE:
OTHER SOURCE(S):

CASREACT 115:183735

GI

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/ Structure 23 in file .gra /
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AB Selective deacetoxylation of O-acetyl sugar lactones, e.g. I (R = OAc), and O-acetyl-3-deoxy sugar lactones, e.g. I (R = H), was accomplished at room temp. with the aid of SmI2 in the presence of Me3CCO2H or AcOH to give the .alpha.,.beta.-unsatd. lactones and satd. lactones (2,3-dideoxy sugar derivs.), resp.

L4 ANSWER 32 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1991:450188 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***115:50188***

TITLE: Novel method for the synthesis of 2',3'-unsaturated

nucleosides from 2'(3')-acetoxy-3'(2')-halogeno derivatives by using sodium dithionite with viologen

as a reductive elimination mediator

AUTHOR(S): Amino, Yusuke; Iwagami, Hisao

CORPORATE SOURCE: Cent. Res. Lab., Ajinomoto Co., Inc., Kawasaki, 210,

Japan

SOURCE: Chemical & Pharmaceutical Bulletin (1991), 39(3),

622-5

CODEN: CPBTAL; ISSN: 0009-2363

DOCUMENT TYPE: Journal LANGUAGE: English

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OTHER SOURCE(S):
                         CASREACT 115:50188
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AB

/ Structure 24 in file .gra / Reductive elimination of vicinal acetylated halohydrins with Na2S2O4 as AB the reducing agent and viologen as the redn. mediator in a 2-phase water-org. system is described. 2',3'-Unsatd. nucleosides such as 1-(5-0-acetyl-2,3-dideoxy-.beta.-D-glycero-pent-2-enofuranosyl)adenine (I), -hypoxanthine, and -uracil were obtained from 2'(3')-acetoxy-3'(2')halogeno derivs., e.g., II, in good yields by means of this procedure. ANSWER 33 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: ***114:194739*** DOCUMENT NUMBER: Electrochemical synthesis of 2',3'-dideoxy-2',3'-TITLE: didehydrocytidine and 2',3'-dideoxycytidine Johansen, Oddvar; Holan, George; Marcuccio, Sebastian AUTHOR(S): M.: Mau, Albert W. H. Div. Chem. Polym., CSIRO, Clayton, 3168, Australia CORPORATE SOURCE: Australian Journal of Chemistry (1991), 44(1), 37-41 SOURCE: CODEN: AJCHAS; ISSN: 0004-9425 DOCUMENT TYPE: Journal English LANGUAGE: CASREACT 114:194739 OTHER SOURCE(S): An improved electrochem. procedure for the synthesis of N4,05-diacetyl-2',3'-dideoxy-2',3'-didehydrocytidine (I) has been established. I was used as an intermediate in the syntheses of 2',3'-dideoxy-2'-3'-didehydrocytidine and 2',3'-dideoxycytidine. ANSWER 34 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN 1991:102667 CAPLUS <<LOGINID::20071011>> ACCESSION NUMBER: ***114:102667*** DOCUMENT NUMBER: The use of acetyl bromide for the multigram synthesis TITLE: of the anti-HIV agent 2',3'-didehydro-2',3'dideoxycytidine (d4C) Starrett, John E., Jr.; Tortolani, David R.; Baker, AUTHOR(S): David C.; Omar, Mahmoud T.; Hebbler, Anna K.; Wos, John A.; Martin, John C.; Mansuri, Muzammil M. Dep. Antiinfect. Chem., Bristol-Myers Squibb Co., CORPORATE SOURCE: Wallingford, CT, 06492, USA SOURCE: Nucleosides & Nucleotides (1990), 9(7), 885-97 CODEN: NUNUD5; ISSN: 0732-8311 DOCUMENT TYPE: Journal English LANGUAGE: OTHER SOURCE(S): CASREACT 114:102667 GI / Structure 25 in file .gra / Treatment of uridine with AcBr produced bromoacetate I, which was reduced

with Zn/Cu to give unsatd. uridine II (R = Ac, X = O). Treatment of II (R = Bz, X = O) with Lawesson's reagent gave thioamide II (R = Bz, X = S) (III). Amination and deprotection of III with NH3/MeOH afforded title

d4C. This multigram scale process for the synthesis of d4C proceeded in 20% yield from uridine.

L4 ANSWER 35 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1990:532652 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***113:132652***

TITLE: Vitamin C and isovitamin C derived chemistry. 4.

Synthesis of some novel furanone chirons

AUTHOR(S): Vekemans, Jozef A. J. M.; Dapperens, Cornelis W. M.;

Claessen, Ron; Koten, Annemiek M. J.; Godefroi, Erik

F.; Chittenden, Gordon J. F.

CORPORATE SOURCE: Dep. Chem. Technol., Univ. Technology, Eindhoven, 5600

MB, Neth.

SOURCE: Journal of Organic Chemistry (1990), 55(19), 5336-44

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 113:132652

GI

/ Structure 26 in file .gra /

The utility of the vitamin C and isovitamin C derived dibromo diacetates I (R = H, R1 = OAc1, R = OAc, R1 = H) for prepg. synthetically useful chirons is further examd. Methodologies for transforming the readily accessible compds. I into 3,6- and 5,6-anhydrohexono-1,4-lactones II (R2 = H, R3 = OH; R2 = OH, R3 = H) and III [(R),(S)-oxirane] are presented. Elaboration of the lactone epoxides III and IV provided optically pure (S,S)-, meso-, and (R,R)-4,4'-bis-.gamma.-butyrolactones V (S,S, meso) and VI. The difference in reactivity between some isomeric intermediates was related to the spatial arrangement at the reactive sites.

L4 ANSWER 36 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1990:441211 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***113:41211***

TITLE: Preparation of pyrrolo[2,3-d]pyrimidine nucleosides as

antiviral agents

INVENTOR(S): Townsend, Leroy B.; Drach, John C.; Krawczyk, Steven

Η.

PATENT ASSIGNEE(S): University of Michigan, USA

SOURCE: U.S., 14 pp.
CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

KIND DATE DATE PATENT NO. APPLICATION NO. --------------_____ US 1987-127487 19900109 19871201 US 4892865 Α PRIORITY APPLN. INFO.: US 1987-127487 19871201

OTHER SOURCE(S): CASREACT 113:41211; MARPAT 113:41211

GI

/ Structure 27 in file .gra /

AB The title compds. [I; R1 = NH2, OH; R2 = H, cyano, CO2H, CONH2, C(S)NH2, etc.; R3 = 2',3'-dideoxy-2',3'-didehydro-.beta.-D-ribofuranosyl,

2',3'-dideoxyribofuranosyl], inhibitors of herpes virus and cytomegalovirus, were prepd. Tubercidin was 3'-brominated-2',5'-di-0acetylated with AcOCMe2COBr in MeOH-MeCN contg. Ac20 and pyridine and the product treated with Zn-Cu in DMF to give, after deacetylation, 2',3'-dideoxy-2',3'-didehydrotubercercidin, whose IC50 for cytomegalovirus was >100 .mu.M. The cytotoxicity of I was also tested.

ANSWER 37 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN L4

ACCESSION NUMBER:

113:41201 DOCUMENT NUMBER:

Synthesis and anti-HIV evaluation of TITLE:

2',3'-dideoxyribo-5-chloropyrimidine analogs: reduced

toxicity of 5-chlorinated 2',3'-dideoxynucleosides

Van Aerschot, Arthur; Everaert, Dirk; Balzarini, Jan;

Augustyns, Koen; Jie, Liu; Janssen, Gerard; Peeters,

Oswald; Blaton, Norbert; De Ranter, Camiel; et al.

Rega Inst. Med. Res., Kathol. Univ. Leuven, Louvain,

B-3000, Belq.

Journal of Medicinal Chemistry (1990), 33(6), 1833-9 SOURCE:

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE: Journal

English LANGUAGE: CASREACT 113:41201

OTHER SOURCE(S):

CORPORATE SOURCE:

AUTHOR(S):

/ Structure 28 in file .gra /

In view of the selective anti-HIV activity of 2',3'-dideoxy-3'-fluoro-5-AB chlorouridine (I), a series of eight 2',3'-dideoxy-5-chloropyrimidines were synthesized and evaluated for their inhibitory activity against human immunodeficiency virus type 1 (HIV-1) replication in MT-4 cells. A marked improvement in selectivity was noted for the 5-chlorouracil derivs. of 2,3-dideoxyribofuranose, 3-azido-2,3-dideoxyribofuranose, and 3-fluoro-2,3-dideoxyribofuranose, mainly due to decreased toxicity of the compds. for the host cells. While chlorination of 2',3'-dideoxycytidine removed the anti-HIV activity, introduction of Cl at C(5) of 3'-fluoro-, 3'-azido- or 2',3'-didehydro-2',3'-dideoxycytidine led to reduced cytotoxicity with only slightly reduced anti-HIV activity. X-ray anal. revealed no close resemblance of I to 3'-azido-3'-deoxythymidine (AZT).

ANSWER 38 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN L4

ACCESSION NUMBER: 1990:119355 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER:

112:119355

Preparation of 2',3'-dideoxy-2',3'-didehydro-TITLE:

> pyrimidine, -azapyrimidine, or -deazapyrimidine nucleosides as antiviral agents against human

immunodeficiency viruses (HIV)

Starrett, John E., Jr.; Mansuri, Muzammil M.; Martin, INVENTOR(S):

John C.; Fuller, Carl E.; Howell, Henry G.

PATENT ASSIGNEE(S): Bristol-Myers Co., USA

Eur. Pat. Appl., 22 pp. SOURCE:

CODEN: EPXXDW

DOCUMENT TYPE: Patent

English LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

DATE DATE APPLICATION NO. PATENT NO. KIND

													_	
ED.	334368			A2		890927	 E1	 -	1989-	1051	269			19890323
	334368			A3		911227		-	1909-	103	209			19090323
EP	R: AT,	BE	СН					ריד	. т.т.	T.II	NT.	SE		
IIS	4904770	55,	O11,	A		900227			1988-			-		19880324
	1339448			C		970909			1989-					19890315
	8901208			A		9890925			1989-					19890320
	171367			В		9921123			1000	1200	,			13030320
	8901338			A		9890925		г	1989-	1225	R			19890321
	93111			В		941115		_	1000	133				17070321
	93111			C		950227								
	105572			A		940412		۲.	1989-	105	572			19890321
	105572			A		940412			1989-					19890321
	105573			A		940530			1989-					19890321
	89693			A		940826			1989-					19890321
	105570			A		940826			1989-					19890321
	8901464			A		890925			1989-					19890322
	8902166			A		920226			1989-					19890322
	8931673			A		890928			1989-					19890323
	622439			B2		920409		-						
	02149595			A		9900608		Р	1989-	715	92			19890323
	140889			B1		980615			1989-					19890323
	5130421			A		920714		5	1991-	697	512			19910429
	9103456			A		890925		2	1991-	345	6			19910903
	172345			В		930329								
	172345			C	19	9930707	,							
	9103457			Α	19	890925	NO	0	1991-	345	7			19910903
ОИ	171314			В	19	921116								
МО	171314			С	19	9930224	:							
NO	9103458			A	19	890925	N	2	1991-	345	8			19910903
NO	171315			В	19	921116	;							
МО	171315			C	19	9930224								
NO	9103459			Α	19	890925	NO.	Э	1991-	345	9			19910903
NO	171316			В	19	9921116	;							
МО	171316			C	19	9930224	:							
US	5212294			Α		9930518		-	1992-					19920331
CA	133,9861			C		9980512			1993-					19931206
FI	9405698			Α		9941202			1994-					19941202
	9405699			Α		9941202			1994-					19941202
	9405700			Α		9941202			1994-					19941202
	1339483			С		9970930			1996-					19960321
	142109			B1		9980701			1997-					19971023
	142110			B1	19	9980701			1997-				_	19971023
PRIORIT	Y APPLN.	INFO	. :						1988-				A	19880324
									1989-					19890315
									1989-					19890320
									1989-					19890321
									1989-				A	19890323
									1989-					19891124
									1991-		91 2			19910429 19940110
							F.	Τ.	1994-	. T 0 3			Α	T2240TT0

OTHER SOURCE(S): CASREACT 112:119355; MARPAT 112:119355

/ Structure 29 in file .gra /

AB 2',3'-Dideoxy-2',3'-didehydronucleosides [I; X = N, CH; Y = CR5; N; Z = CH, N; R4 = OH, NH2; R5 = H, (halo-substituted) CnH2nA or (CH2)mCH:CHA; m

= 0-3; n = 1-3; A = H, F, C1, Br, iodo], useful as antiviral agents, esp. against HIV, are prepd. in high yields and on a relatively large scale by subjecting various intermediates II-IV, V (R = Br, R1 = isobutyryloxy; or R = isobutyryloxy, R1 = Br) and VI to elimination reactions by treatment of (1) II with a strong base (e.g. tert-BuOK), (2) II with an org. acid in Ac20 at 120-160.degree. for 4-8 h followed by 5-0-deacetylation, (3) IV with P(OEt)3 in a polar solvent at 140-175.degree. 0.5-4 h, (4) V with Zn/Cu in an aprotic solvent, and (5) VI with a nonnucleophilic base (e.g. Bu4NF) or nucleophilic base (e.g. tert-BuOK and KOH). Thus, mesylation of thymidine with MeSO2Cl in pyridine at 0-5.degree. gave 81% 3',5'-di-O-(methanesulfonyl)thymidine which was added portionwise to a stirred soln. of aq. NaOH and then refluxed 2 h to give 74% 1-(3,5-anhydro-2-deoxy-.beta.-D-threo-pentofuranosyl)thymine. stirred soln. of 90.0 g of the latter octane was added 97% tert-BuOK (74 q) portionwise over 25 min at 18-22.degree. in an ice bath. The mixt. was stirred 1 h to give 57% 1-(2,3-dideoxy-.beta.-glycero-pent-2enofuranosyl)thymine (VII). VII showed an IC50 of 0.33 .mu.M against HIV in CEM cells vs. 0.45 for AZT.

L4 ANSWER 39 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1989:554286 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***111:154286***

TITLE: Preparation of 1-(2,3-dideoxy-.beta.-D-glycero-pent-2-

enofuranosyl)thymine (d4T) and 2',3'-dideoxyadenosine

(ddA): general methods for the synthesis of

2',3'-olefinic and 2',3'-dideoxy nucleoside analogs

active against HIV

AUTHOR(S): Mansuri, Muzammil M.; Starrett, John E., Jr.; Wos,

John A.; Tortolani, David R.; Brodfuehrer, Paul R.;

Howell, Henry G.; Martin, John C.

CORPORATE SOURCE: Pharm. Res. Dev. Div., Bristol-Myers, Wallingford, CT,

06492-7660, USA

SOURCE: Journal of Organic Chemistry (1989), 54(20), 4780-5

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 111:154286

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/ Structure 30 in file .gra /

AB Methods for the prepn. of the 2',3'-unsatd. thymidine and cytidine analogs (I; R = Me, R1 = OH; R = H, R1 = NH2), 2',3'-dideoxycytidine and 2',3'-dideoxyadenosine, which are active in vitro against HIV, are reported. The methods used were the Corey-Winter reaction involving the fragmentation of a cyclic thionocarbonate II, olefin formation from 2',3'-O-alkoxymethylidene cyclic ortho esters, and the reductive elimination of the 2',3' halo acetates, e.g., III [R2 = COC(OAc)Me2, R3 = H, Me; R1 = Ac, R3 = Me). Of these 3 methods, the last was the most versatile, since the intermediates III or the trans-3'(2')-bromo-2'(3')-O-acetyl-3'(2')-deoxyarabinosylpurines are readily transformed to the corresponding olefins. As an example of the prepn. of a satd. 2',3'-dideoxy analog, 2',3'-dideoxyadenosine was obtained by catalytic redn. of the corresponding olefinic nucleoside.

L4 ANSWER 40 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1989:439794 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***111:39794***

2',3'-Dideoxyadenosine analogs of the nucleoside TITLE: antibiotics tubercidin, toyocamycin and sangivamycin Krawczyk, Steven H.; Townsend, Leroy B. AUTHOR(S): Coll. Pharm., Univ. Michigan, Ann Arbor, MI, CORPORATE SOURCE: 48109-1065, USA Nucleosides & Nucleotides (1989), 8(1), 97-115 SOURCE: CODEN: NUNUD5; ISSN: 0732-8311

Journal DOCUMENT TYPE: LANGUAGE: English

OTHER SOURCE(S): CASREACT 111:39794

/ Structure 31 in file .gra /

Dideoxynucleosides I (R1 = H, cyano, CONH2) were prepd. Toyocamycin was AB treated with Me2C(OAc)COBr, MeOH, and Ac2O to give bromodeoxy compd. II, and II was subjected to sequential elimination reaction with Zn-Cu, deacetylation, and hydrogenation to give I (R1 = cyano).

ANSWER 41 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

1988:75761 CAPLUS <<LOGINID::20071011>> ACCESSION NUMBER:

DOCUMENT NUMBER: ***108:75761***

Vitamin C and isovitamin C derived chemistry. 3. TITLE:

Chiral butenolides via efficient 2,3-

didehydroxylations of L-qulono-, D-mannono-, and

D-ribono-1,4-lactones

Vekemans, Jozef A. J. M.; Franken, Gabriel A. M.; AUTHOR(S):

Dapperens, Cornelis W. M.; Godefroi, Erik F.;

Chittenden, Gordon J. F.

Dep. Chem. Technol., Univ. Technol., Eindhoven, 5600 CORPORATE SOURCE:

MB, Neth.

Journal of Organic Chemistry (1988), 53(3), 627-33 SOURCE:

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal English LANGUAGE:

OTHER SOURCE(S): CASREACT 108:75761

/ Structure 32 in file .gra /

Efficient, operationally simple procedures for prepg. the chiral AB butenolides I (R = H, R1 = R2 = OH; R = R2 = OH, R1 = H; R = H, R1 = OAc,R2 = Br; R = OAc, R1 = H, R2 = Br; R = H, R1 = OH, R2 = Br; R = OH, R1 = H, R2 = Br; R = H, R1R2 = O; RR2 = O, R1 = H) from com. L-ascorbic acid and D-isoascorbic acid are described. The concept centers on the novel NaHSO3-induced regiospecific trans-.beta.-bromo-acetoxy elimination of the readily accessible O-acetylated bromodeoxyaldono-1,4-lactones II (R = H, R1 = OAc; R = OAc, R1 = H) to I (R2 = Br). These, on deacetylation and treatment of the resulting bromohydrins with Ag2O, afford the enantiomerically pure epoxides I (R = H, R1R2 = O; RR2 = O, R1 = H) and thence, in boiling water, the corresponding diols I (R = H, R1 = R2 = OH; R = R2 = OH, R1 = H). Lactone III was also prepd. in 48% overall yield from D-ribono-1,4-lactone.

ANSWER 42 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER:

DOCUMENT NUMBER: ***108:56502***

TITLE: An efficient synthesis of (S)-5-hydroxymethyl-2(5H)-

furanone

AUTHOR(S): Vekemans, Jozef A. J. M.; Franken, Gabriel A. M.;

Chittenden, Gordon J. F.; Godefroi, Erik F.

CORPORATE SOURCE: Dep. Chem. Technol., Univ. Technol., Eindhoven, 5600

MB, Neth.

SOURCE: Tetrahedron Letters (1987), 28(20), 2299-300

CODEN: TELEAY; ISSN: 0040-4039

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 108:56502

CT

/ Structure 33 in file .gra /

AB A three step synthesis of the title compd. [I; R = OH (II)] from D-ribono-1,4-lactone in 48% yield is described. The method centers on a novel NaHSO3-induced trans-Br-OAc elimination of the readily prepd. bromoacetate mixt. III (R1 = OAc, Br) to the corresponding butenolides I (R = OAc, Br), which (R = OAc) on hydrolysis and purifn., yields pure II.

L4 ANSWER 43 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1987:196718 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***106:196718***

TITLE: The reaction of 9-(2',3'-dideoxy-.beta.-D-glycero-pent-

2-enofuranosyl)adenine derivatives with arene- and alkanesulfenyl chlorides. An unusual ring opening

reaction of thiiranium ions

AUTHOR(S): Welch, C. J.; Bazin, H.; Chattopadhyaya, J.

CORPORATE SOURCE: Biomed. Cent., Uppsala Univ., Uppsala, S-751 23, Swed.

SOURCE: Acta Chemica Scandinavica, Series B: Organic

Chemistry and Biochemistry (1986), B40(5), 343-57

CODEN: ACBOCV; ISSN: 0302-4369

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 106:196718

GT

/ Structure 34 in file .gra /

AB A series of arenesulfenyl chlorides and methanesulfenyl chloride were treated in pyridine soln. with the title adenine deriv. I. In each case, the relative abundance of isomeric .beta.-chloroarene/alkane sulfides was estd. and used to assess the relative thermodn. stabilities of the lyxo and ribo thiiranium ion intermediates and their steric susceptibilities to attack by a nucleophile (Cl-). Subsequently, the .beta.-chloroarene/alkane sulfides II [R = Cl, Rl = R2 = H, R3 = Q (Q = PhS, p-MeC6H4S, o-MeC6H4S, p-ClC6H4S, MeS); R = R3 = H, R1 = Cl, R2 = Q; R = Q, R1 = R2 = H, R3 = Cl; R = R3 = H, R1 = Q, R2 = Cl], upon alk. treatment, underwent an unprecedented ylide-mediated .beta.-elimination of the intermediary lyxo and ribo thiiranium ions, to give only 2'-ene-thiol ethers III (R4 = Q).

L4 ANSWER 44 OF 44 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1985:422870 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***103:22870***

TITLE: Elimination reaction of aldehydo-D-glucurono-6,3-

lactone derivatives

Itoh, Hiroshi; Aoki, Koichi; Matsumoto, Takahiro AUTHOR(S):

Noguchi Inst., Tokyo, 173, Japan CORPORATE SOURCE:

SOURCE: Bulletin of the Chemical Society of Japan (1985),

58(2), 777-8

CODEN: BCSJA8; ISSN: 0009-2673

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 103:22870

AB D-Glucurono-6,3-lactone derivs. are converted to butenolide or

dihydro-2H-pyran derivs. by treatment with Et3N.

=> file stnq

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

124.99 239.20

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE TOTAL ENTRY SESSION

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0.12 239.32 FULL ESTIMATED COST

SINCE FILE TOTAL DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

ENTRY SESSION

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L1

(FILE 'HOME' ENTERED AT 15:31:21 ON 11 OCT 2007)

FILE 'CASREACT' ENTERED AT 15:31:37 ON 11 OCT 2007

STRUCTURE UPLOADED

L2 4 S L1

L3 44 S L1 FULL

FILE 'CAPLUS' ENTERED AT 15:32:41 ON 11 OCT 2007

L4 44 S L3

FILE 'STNGUIDE' ENTERED AT 15:33:12 ON 11 OCT 2007

FILE 'CAPLUS' ENTERED AT 15:34:30 ON 11 OCT 2007

=> s 14 and zinc

641794 ZINC

139 ZINCS

641819 ZINC

(ZINC OR ZINCS)

L5 5 L4 AND ZINC

=> d 15 1-5 ibib abs

L5 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:1094206 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***143:347399***

TITLE: Method for preparing 2',3'-dideoxycytidine

INVENTOR(S): Lu, Jinkang; Wang, Xinhua

PATENT ASSIGNEE(S): Peop. Rep. China

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 8 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent

LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	AP	PLICATION NO.	DATE		
CN 1563030	A	20050112	CN	2004-10014350	20040315		
PRIORITY APPLN. INFO.:			CN	2004-10014350	20040315		
OFFICE GOLLD OF (G)	CA CDES	OM 142.2472	00				

OTHER SOURCE(S): CASREACT 143:347399

The title method comprises allowing cytidine to react with acetic anhydride to get 4-N-acetylcytidine, reacting 4-N-acetylcytidine with a satd. acetic acid soln. of hydrogen bromide with acetic anhydride as catalyst to get a bromide mixt.; reacting to get 4-N-acetyl-2',3'-didehydro-2',3'-dideoxycytidine-5'-acetate with ***zinc*** copper couple as catalyst; hydrogenating in the presence of Pd/C to get 4-N-acetyl-2',3'-dideoxyguanosine-5'-acetate; and reacting at the presence of methanol and triethylamine to get 2',3'-dideoxycytidine. The invention has the advantages of simple technique, easy operation, easily-accessible raw material, and high yield exceeding 90%.

L5 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003:123466 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***138:137526***

TITLE: Preparation of 2',3'-didehydro-3'-deoxythymidine

INVENTOR(S): Yao, Qizheng; Lu, Gang

PATENT ASSIGNEE(S): China Pharmacy Univ., Peop. Rep. China

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 12 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

CN 1324800 A 20011205 CN 2001-113661 20010601

PRIORITY APPLN. INFO.: CN 2001-113661 20010601

OTHER SOURCE(S): CASREACT 138:137526; MARPAT 138:137526

Stavudine (2',3'-didehydro-3'-deoxythymidine) as antiviral agent is synthesized by dehydrating 5-methyluridine with di-C1- 4 alkyl or di-Ph carbonate (at a molar ratio of 1.1-1.5) in anhyd. polar solvent (such as DMF or DMA) in the presence of base (such as NaHCO3 or KHCO3) at 90-150.degree. to obtain 2,2'-anhydro-5-methyluridine; acylating and halogenating with C1- 6 acyl halide (or org. anhydride-org. acid-NaBr and/or H2SO4, or org. anhydride-pyridinium bromide perbromide) in polar solvent (such as acetonitrile, DMF, or Et acetate) at 50-75.degree. to obtain 2'-halo-3',5'-O-diacylthymidine (halo = C1, Br, or I); reductive eliminating with metal reductant (such as Zn or Zn-Cu) in methanol or ethanol and/or Et acetate in the presence of acid (such as acetic acid or propanoic acid) at 5-35.degree.; and hydrolyzing with base (such as Na methoxide or NH3) in methanol or ethanol and/or dioxane.

L5 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:762991 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***135:304107***

TITLE: Process for preparing 5'-acetylstavudine by the

reductive elimination reaction of 2'-deoxy-2'-bromo-

3',5'-diacetyl-5-methyluridine

INVENTOR(S): Bertolini, Giorgio; Frigerio, Marco; Velati, Maurizio;

Petrucciani, Luigi

PATENT ASSIGNEE(S): Clariant Life Science Molecules, Italy

SOURCE: PCT Int. Appl., 12 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAC	rent :	NO.			KIN	D	DATE		API	PLICAT	Ι	DATE				
WO					A1 20011018			1018	WO 2001-IB588					20010410		
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		PT,	SE,	TR												
IT	IT 2000MI0810						20011012 IT 2000-MI810							200004	12	
. IT	IT 1318463						2003	0825								
EP	1274	704			A1		2003	0115	EP	2001-	917385		2	200104	10	
EP	EP 1274704						2004	0114								
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ES	2213	109			Т3		2004	0816	ES	2001-	191738	5	2	200104	10	
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PRIORITY	PRIORITY APPLN. INFO.:								IT	2000-	MI810		A 2	200004	12	
									WO	2001-	IB588		W 2	200104	10	

OTHER SOURCE(S): CASREACT 135:304107

AB 5'-Acetylstavudine is prepd. in high yield and selectivity by the reaction of 2'-deoxy-2'-bromo-3',5'-diacetyl-5-methyluridine in the presence of ***zinc*** as reducing agent combined with an activating agent which is an ammonium salt (e.g., triethylammonium hydrochloride) and/or a phosphonium salt in a dipolar aprotic solvent (e.g., DMF).

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS

L5 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1993:496031 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***119:96031***

TITLE: An efficient and general synthesis of 5'-esters of

2',3'-didehydro-2',3'-dideoxynucleosides: a facile opening of 2',3'-orthoacetates of ribonucleosides followed by reductive elimination of the halo acetates

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Talekar, Ratnakar R.; Coe, Paul L.; Walker, Richard T.

CORPORATE SOURCE: Sch. Chem., Univ. Birmingham, Birmingham, B15 2TT, UK

SOURCE: Synthesis (1993), (3), 303-6

CODEN: SYNTBF; ISSN: 0039-7881

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 119:96031

OTHER SOURC

AUTHOR(S):

GI

/ Structure 35 in file .gra /

AB A three-step reaction sequence starting from a ribonucleoside I [B = N4-acetylcytosinyl (C), uracilyl (U), 6-oxopurinyl (I), adeninyl (A)] to give the corresponding 5'-O-acyl-2',3'-didehydro-2',3'-dideoxynucleoside II is described. The key intermediate is the bromoacetate III (B = C, I, A, R1 = Br, R2 = H, R3 = OAc; B = U, R1 = H, R2 = OAc, R3 = Br), made by reaction of the 2',3'-methoxyethylidene nucleoside IV with acetyl bromide. Reductive elimination of the bromoacetate using a ***zinc*** -copper couple furnishes the desired compds. in good overall yield.

L5 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1987:196718 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***106:196718***

TITLE: The reaction of 9-(2',3'-dideoxy-.beta.-D-glycero-pent-

2-enofuranosyl) adenine derivatives with arene- and alkanesulfenyl chlorides. An unusual ring opening

reaction of thiiranium ions

AUTHOR(S): Welch, C. J.; Bazin, H.; Chattopadhyaya, J.

CORPORATE SOURCE: Biomed. Cent., Uppsala Univ., Uppsala, S-751 23, Swed.

SOURCE: Acta Chemica Scandinavica, Series B: Organic

Chemistry and Biochemistry (1986), B40(5), 343-57

CODEN: ACBOCV; ISSN: 0302-4369

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 106:196718

GI

AB A series of arenesulfenyl chlorides and methanesulfenyl chloride were treated in pyridine soln. with the title adenine deriv. I. In each case, the relative abundance of isomeric .beta.-chloroarene/alkane sulfides was estd. and used to assess the relative thermodn. stabilities of the lyxo and ribo thiiranium ion intermediates and their steric susceptibilities to attack by a nucleophile (Cl-). Subsequently, the .beta.-chloroarene/alkane sulfides II [R = Cl, Rl = R2 = H, R3 = Q (Q = PhS, p-MeC6H4S, o-MeC6H4S, p-ClC6H4S, MeS); R = R3 = H, R1 = Cl, R2 = Q; R = Q, R1 = R2 = H, R3 = Cl; R = R3 = H, R1 = Q, R2 = Cl], upon alk. treatment, underwent an unprecedented ylide-mediated .beta.-elimination of the intermediary lyxo and ribo thiiranium ions, to give only 2'-ene-thiol ethers III (R4 = Q).

=> FIL STNGUIDE

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 16.63 255.95

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE TOTAL
ENTRY SESSION

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L4

L5

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FILE 'CASREACT' ENTERED AT 15:31:37 ON 11 OCT 2007

L1 STRUCTURE UPLOADED

L2 4 S L1

L3 44 S L1 FULL

FILE 'CAPLUS' ENTERED AT 15:32:41 ON 11 OCT 2007 44 S L3

FILE 'STNGUIDE' ENTERED AT 15:33:12 ON 11 OCT 2007

FILE 'CAPLUS' ENTERED AT 15:34:30 ON 11 OCT 2007
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FILE 'STNGUIDE' ENTERED AT 15:34:52 ON 11 OCT 2007

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L5 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:1094206 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***143:347399***

TITLE: Method for preparing 2',3'-dideoxycytidine

INVENTOR(S): Lu, Jinkang; Wang, Xinhua

PATENT ASSIGNEE(S): Peop. Rep. China

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 8 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

OTHER SOURCE(S): CASREACT 143:347399
AN 2005:1094206 CAPLUS <<LOGINID::20071011>>

DN ***143:347399***

AB

. . . acid soln. of hydrogen bromide with acetic anhydride as catalyst to get a bromide mixt.; reacting to get 4-N-acetyl-2',3'-didehydro-2',3'-dideoxycytidine-5'-acetate with ***zinc*** copper couple as catalyst; hydrogenating in the presence of Pd/C to get 4-N-acetyl-2',3'-dideoxyguanosine-5'-acetate; and reacting at the presence of methanol. .

5 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003:123466 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***138:137526***

TITLE: Preparation of 2',3'-didehydro-3'-deoxythymidine

INVENTOR(S): Yao, Qizheng; Lu, Gang

PATENT ASSIGNEE(S): China Pharmacy Univ., Peop. Rep. China

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 12 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

CN 1324800 A 20011205 CN 2001-113661 20010601

PRIORITY APPLN. INFO.: CN 2001-113661 20010601

OTHER SOURCE(S): CASREACT 138:137526; MARPAT 138:137526

AN 2003:123466 CAPLUS <<LOGINID::20071011>>

DN ***138:137526***

IT 598-22-1, Propionyl bromide 1463-10-1 7440-66-6, ***Zinc*** , reactions 12621-78-2

RL: RCT (Reactant); RACT (Reactant or reagent)
 (prepn. of 2',3'-didehydro-3'-deoxythymidine)

L5 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:762991 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***135:304107***

TITLE: Process for preparing 5'-acetylstavudine by the reductive elimination reaction of 2'-deoxy-2'-bromo-

3',5'-diacetyl-5-methyluridine

INVENTOR(S): Bertolini, Giorgio; Frigerio, Marco; Velati, Maurizio;

Petrucciani, Luigi

PATENT ASSIGNEE(S): Clariant Life Science Molecules, Italy

SOURCE: PCT Int. Appl., 12 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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PRIORITY APPLN. INFO.:
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                                          WO 2001-IB588
                                                            W 20010410
                        CASREACT 135:304107
OTHER SOURCE(S):
                             THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                              RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
    AN
DN
      ***135:304107***
    5'-Acetylstavudine is prepd. in high yield and selectivity by the reaction
AB
    of 2'-deoxy-2'-bromo-3',5'-diacetyl-5-methyluridine in the presence of
                   as reducing agent combined with an activating agent which is
    an ammonium salt (e.g., triethylammonium hydrochloride) and/or a
    phosphonium salt.
    7440-66-6, ***Zinc***
                            , reactions
IT
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (powder; process for prepg. 5'-acetylstavudine by the reductive
       elimination reaction of 2'-deoxy-2'-bromo-3',5'-diacetyl-5-
       methyluridine using)
    ANSWER 4 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN
                        1993:496031 CAPLUS <<LOGINID::20071011>>
ACCESSION NUMBER:
DOCUMENT NUMBER:
                          ***119:96031***
                        An efficient and general synthesis of 5'-esters of
TITLE:
                        2',3'-didehydro-2',3'-dideoxynucleosides: a facile
                        opening of 2',3'-orthoacetates of ribonucleosides
                        followed by reductive elimination of the halo acetates
                        Talekar, Ratnakar R.; Coe, Paul L.; Walker, Richard T.
AUTHOR(S):
                        Sch. Chem., Univ. Birmingham, Birmingham, B15 2TT, UK
CORPORATE SOURCE:
                        Synthesis (1993), (3), 303-6
SOURCE:
                        CODEN: SYNTBF; ISSN: 0039-7881
DOCUMENT TYPE:
                        Journal
                        English
LANGUAGE:
OTHER SOURCE(S):
                        CASREACT 119:96031
    1993:496031 CAPLUS <<LOGINID::20071011>>
AN
      ***119:96031***
DN
     . . . = Br), made by reaction of the 2',3'-methoxyethylidene nucleoside
AB
    IV with acetyl bromide. Reductive elimination of the bromoacetate using a
      ***zinc*** -copper couple furnishes the desired compds. in good overall
    yield.
    didehydrodideoxynucleoside; nucleoside didehydrodideoxy; ribonucleoside
ST
    reaction orthoformate; methoxyethylidene nucleoside reaction acetyl
    bromide; bromodeoxy nucleoside redn ***zinc*** copper couple
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20011018

Α1

WO 2001-IB588

20010410

ACCESSION NUMBER: DOCUMENT NUMBER: ***106:196718***

WO 2001077103

1987:196718 CAPLUS <<LOGINID::20071011>>

ANSWER 5 OF 5 CAPLUS COPYRIGHT 2007 ACS on STN

TITLE: The reaction of 9-(2',3'-dideoxy-.beta.-D-glycero-pent-

2-enofuranosyl) adenine derivatives with arene- and alkanesulfenyl chlorides. An unusual ring opening

reaction of thiiranium ions

AUTHOR(S): Welch, C. J.; Bazin, H.; Chattopadhyaya, J.

CORPORATE SOURCE: Biomed. Cent., Uppsala Univ., Uppsala, S-751 23, Swed.

SOURCE: Acta Chemica Scandinavica, Series B: Organic

Chemistry and Biochemistry (1986), B40(5), 343-57

108183-22-8P 108183-23-9P 108194-70-3P 108194-71-4P

CODEN: ACBOCV; ISSN: 0302-4369

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 106:196718
AN 1987:196718 CAPLUS <<LOGINID::20071011>>

DN ***106:196718***

IT

64-19-7P, preparation 67-56-1P, preparation 64-17-5P, preparation 108-24-7P 109-86-4P, 2-Methoxyethanol 109-99-9P, preparation 110-86-1P, Pyridine, preparation 124-41-4P 127-09-3P 7440-02-0P, Nickel, preparation 7440-66-6P, ***Zinc*** , preparation 7664-41-7P, preparation 7722-64-7P 7732-18-5P, Water, preparation 104525-44-2P 104525-45-3P 104525-46-4P 104525-47-5P 104525-48-6P 104525-50-0P 104525-51-1P 108182-98-5P 108182-99-6P 104525-49-7P 108183-00-2P 108183-01-3P 108183-07-9P 108183-09-1P 108183-10-4P 108183-11-5P 108183-12-6P 108183-13-7P 108183-14-8P 108183-20-6P

108183-21-7P 108194-72-5P

=> file stng

COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST
0.06
268.75

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE TOTAL

ENTRY SESSION

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DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE TOTAL

ENTRY SESSION

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FILE COVERS 1907 - 11 Oct 2007 VOL 147 ISS 16 FILE LAST UPDATED: 10 Oct 2007 (20071010/ED)

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=> d his

L1

L5

(FILE 'HOME' ENTERED AT 15:31:21 ON 11 OCT 2007)

FILE 'CASREACT' ENTERED AT 15:31:37 ON 11 OCT 2007

STRUCTURE UPLOADED

L2 4 S L1

L3 44 S L1 FULL

FILE 'CAPLUS' ENTERED AT 15:32:41 ON 11 OCT 2007

L4 44 S L3

FILE 'STNGUIDE' ENTERED AT 15:33:12 ON 11 OCT 2007

FILE 'CAPLUS' ENTERED AT 15:34:30 ON 11 OCT 2007

5 S L4 AND ZINC

FILE 'STNGUIDE' ENTERED AT 15:34:52 ON 11 OCT 2007

FILE 'CAPLUS' ENTERED AT 15:35:17 ON 11 OCT 2007

FILE 'STNGUIDE' ENTERED AT 15:35:17 ON 11 OCT 2007

FILE 'STNGUIDE' ENTERED AT 15:35:22 ON 11 OCT 2007

FILE 'CAPLUS' ENTERED AT 15:36:05 ON 11 OCT 2007

=> s 15 and sulfide

332880 SULFIDE

87975 SULFIDES

366071 SULFIDE

(SULFIDE OR SULFIDES)

L6 1 L5 AND SULFIDE

=> d l6 ibib kwic

L6 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1987:196718 CAPLUS <<LOGINID::20071011>>

DOCUMENT NUMBER: ***106:196718***

TITLE: The reaction of 9-(2',3'-dideoxy-.beta.-D-glycero-pent-

2-enofuranosyl) adenine derivatives with arene- and alkanesulfenyl chlorides. An unusual ring opening

reaction of thiiranium ions

AUTHOR(S): Welch, C. J.; Bazin, H.; Chattopadhyaya, J.

CORPORATE SOURCE: Biomed. Cent., Uppsala Univ., Uppsala, S-751 23, Swed.

SOURCE: Acta Chemica Scandinavica, Series B: Organic

```
Chemistry and Biochemistry (1986), B40(5), 343-57
                        CODEN: ACBOCV; ISSN: 0302-4369
DOCUMENT TYPE:
                        Journal
LANGUAGE:
                       English
                       CASREACT 106:196718
OTHER SOURCE(S):
    ***106:196718***
DN
     . . . were treated in pyridine soln. with the title adenine deriv. I.
AB
    In each case, the relative abundance of isomeric .beta.-chloroarene/alkane
      ***sulfides*** was estd. and used to assess the relative thermodn.
    stabilities of the lyxo and ribo thiiranium ion intermediates and their
    steric susceptibilities to attack by a nucleophile (Cl-). Subsequently,
    the .beta.-chloroarene/alkane ***sulfides*** II [R = Cl, Rl = R2 = H,
    R3 = Q (Q = PhS, p-MeC6H4S, o-MeC6H4S, p-ClC6H4S, MeS); R. . .
    64-17-5P, preparation 64-19-7P, preparation 67-56-1P, preparation
IT
    98-88-4P 108-24-7P 109-86-4P, 2-Methoxyethanol 109-99-9P,
                  110-86-1P, Pyridine, preparation 124-41-4P
                                                             127-09-3P
    preparation
    7440-02-0P, Nickel, preparation 7440-66-6P,
                                                  ***Zinc*** , preparation
     7664-41-7P, preparation 7722-64-7P 7732-18-5P, Water, preparation
                   104525-45-3P 104525-46-4P 104525-47-5P
                                                              104525-48-6P
     104525-44-2P
     104525-49-7P 104525-50-0P 104525-51-1P 108182-98-5P
                                                              108182-99-6P
     108183-00-2P 108183-01-3P 108183-07-9P 108183-09-1P 108183-10-4P
     108183-11-5P 108183-12-6P 108183-13-7P 108183-14-8P
                                                              108183-20-6P
     108183-21-7P 108183-22-8P 108183-23-9P 108194-70-3P 108194-71-4P
     108194-72-5P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of)
=> d his
     (FILE 'HOME' ENTERED AT 15:31:21 ON 11 OCT 2007)
     FILE 'CASREACT' ENTERED AT 15:31:37 ON 11 OCT 2007
               STRUCTURE UPLOADED
L1
             4 S L1
L2
            44 S L1 FULL
L3
     FILE 'CAPLUS' ENTERED AT 15:32:41 ON 11 OCT 2007
            44 S L3
L4
     FILE 'STNGUIDE' ENTERED AT 15:33:12 ON 11 OCT 2007
     FILE 'CAPLUS' ENTERED AT 15:34:30 ON 11 OCT 2007
             5 S L4 AND ZINC
L5
     FILE 'STNGUIDE' ENTERED AT 15:34:52 ON 11 OCT 2007
     FILE 'CAPLUS' ENTERED AT 15:35:17 ON 11 OCT 2007
     FILE 'STNGUIDE' ENTERED AT 15:35:17 ON 11 OCT 2007
     FILE 'STNGUIDE' ENTERED AT 15:35:22 ON 11 OCT 2007
     FILE 'CAPLUS' ENTERED AT 15:36:05 ON 11 OCT 2007
             1 S L5 AND SULFIDE
L6
```

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=> LOG Y

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
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CA SUBSCRIBER PRICE	-0.78	-41.34

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     2 JUL 02 LMEDLINE coverage updated
NEWS
NEWS 3 JUL 02 SCISEARCH enhanced with complete author names
NEWS 4 JUL 02 CHEMCATS accession numbers revised
NEWS 5 JUL 02 CA/CAplus enhanced with utility model patents from China
NEWS 6 JUL 16 CAplus enhanced with French and German abstracts
     7 JUL 18 CA/CAplus patent coverage enhanced
NEWS
     8 JUL 26 USPATFULL/USPAT2 enhanced with IPC reclassification
NEWS
NEWS 9 JUL 30 USGENE now available on STN
NEWS 10 AUG 06 CAS REGISTRY enhanced with new experimental property tags
NEWS 11 AUG 06 BEILSTEIN updated with new compounds
NEWS 12 AUG 06 FSTA enhanced with new thesaurus edition
NEWS 13 AUG 13 CA/Caplus enhanced with additional kind codes for granted
                 patents
NEWS 14 AUG 20 CA/CAplus enhanced with CAS indexing in pre-1907 records
NEWS 15 AUG 27
                Full-text patent databases enhanced with predefined
                 patent family display formats from INPADOCDB
        AUG 27
NEWS 16
                USPATOLD now available on STN
NEWS 17
        AUG 28
                CAS REGISTRY enhanced with additional experimental
                 spectral property data
NEWS 18 SEP 07
                 STN AnaVist, Version 2.0, now available with Derwent
                 World Patents Index
        SEP 13 FORIS renamed to SOFIS
NEWS 19
NEWS 20 SEP 13
                INPADOCDB enhanced with monthly SDI frequency
NEWS 21 SEP 17
                CA/CAplus enhanced with printed CA page images from
                 1967-1998
NEWS 22
         SEP 17
                CAplus coverage extended to include traditional medicine
                 patents
         SEP 24
                EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS · 23
NEWS 24
         OCT 02
                CA/CAplus enhanced with pre-1907 records from Chemisches
                 Zentralblatt
NEWS EXPRESS 19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2,
              CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.
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=> file caplus
COST IN U.S. DOLLARS

SINCE FILE TOTAL
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0.21 0.21

FULL ESTIMATED COST

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=> zinc (S) "sodium sulfide"

641794 ZINC

139 ZINCS

641819 ZINC

(ZINC OR ZINCS)

1147007 "SODIUM"

40 "SODIUMS"

1147016 "SODIUM"

("SODIUM" OR "SODIUMS")

332880 "SULFIDE"

87975 "SULFIDES"

366071 "SULFIDE"

("SULFIDE" OR "SULFIDES")

13224 "SODIUM SULFIDE"

("SODIUM"(W) "SULFIDE")

```
=> d scan
```

L1 373 ANSWERS CAPLUS COPYRIGHT 2007 ACS on STN

IC ICM A62D003-00

INCL 588255000; 588252000; 588256000; 044271000; 264004300; 264004700

CC 60-3 (Waste Treatment and Disposal)
Section cross-reference(s): 35, 46, 61

TI Process for disposing of solvent-containing liquids

ST absorbent blend treatment disposal solvent waste

IT Amides, uses

RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses)
(N-(hydroxyalkyl), as solvent type, absorption of; disposal of
solvent-containing liqs. without polluting groundwater by mixing them with
inventive absorbent blend)

IT Wastewater treatment

(absorption; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend)

IT Sulfonic acids, uses

RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses)
(alkanesulfonic, salts, as solvent type, absorption of; disposal of
solvent-containing liqs. without polluting groundwater by mixing them with
inventive absorbent blend)

IT Amine oxides

RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses)
(as solvent type, absorption of; disposal of solvent-containing liqs.
without polluting groundwater by mixing them with inventive absorbent blend)

IT Wastewater treatment

(biol.; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend)

IT Amides, uses

RL: NUU (Other use, unclassified); USES (Uses)
(coco, N,N-bis(hydroxyethyl), Carsamide CA, as solvent, absorption of;
disposal of solvent-containing liqs. without polluting groundwater by
mixing them with inventive absorbent blend)

IT Amides, uses

RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses) (coco, N-[3-(dimethylamino)propyl], N-oxides, Jordamox CAPA, as solvent, absorption of; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend)

IT Absorbents

Crosslinking

Solvents

(disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend)

IT Perlite

Pumice

RL: NUU (Other use, unclassified); USES (Uses)
(disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend)

IT Wastewater treatment

(enzymic; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend)

IT Alcohols, uses

RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses)

(ethoxylated, as solvent type, absorption of; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend) Bacilli (for biodegrdn. of hydrocarbon solvents; disposal of solvent-containing ligs. without polluting groundwater by mixing them with inventive absorbent blend) Enzymes, processes RL: CPS (Chemical process); PEP (Physical, engineering or chemical process); PROC (Process) (for degradation of hydrocarbon solvents; disposal of solvent-containing without polluting groundwater by mixing them with inventive absorbent blend) Solid wastes (formation of; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend) (ground; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend) Groundwater pollution (prevention of; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend) Wastewater treatment (solidification; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend) Wastewater (solvent-containing; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend) (solvents; disposal of solvent-containing ligs. without polluting groundwater by mixing them with inventive absorbent blend) Solvents (waste; disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend) 7440-44-0, Carbon, uses RL: NUU (Other use, unclassified); USES (Uses) (activated; disposal of solvent-containing ligs. without polluting groundwater by mixing them with inventive absorbent blend) 9003-11-6, Ethylene oxide propylene oxide copolymer RL: PEP (Physical, engineering or chemical process); PYP (Physical process); PROC (Process) (as solvent type, absorption of; disposal of solvent-containing ligs. without polluting groundwater by mixing them with inventive absorbent blend) .9002-93-1, Triton X 45 9016-45-9, Polyethylene glycol nonylphenyl ether 25155-30-0, Calsoft F 90 691397-13-4, Pluronic L 61 RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses) (as solvent, absorption of; disposal of solvent-containing ligs. without polluting groundwater by mixing them with inventive absorbent blend) 7439-92-1, Lead, reactions 7439-96-5, Manganese, reactions Mercury, reactions 7440-02-0, Nickel, reactions 7440-31-5, Tin, 7440-33-7, Tungsten, reactions reactions 7440-62-2, Vanadium, 7440-66-6, **Zinc**, reactions reactions

solvent-containing liqs. without polluting groundwater by mixing them with

RL: RCT (Reactant); RACT (Reactant or reagent)

(complexation of, by sodium sulfide; disposal of

ΙT

IT

liqs.

IT

inventive absorbent blend) 1305-78-8, Calcium oxide, uses IT 1344-28-1, Aluminum oxide, uses 7631-86-9, Silicon oxide, uses 7778-18-9, Calcium sulfate 9003-06-9D, 2-Propenamide, polymer with 2-propenoic acid, salts RL: NUU (Other use, unclassified); USES (Uses) (disposal of solvent-containing liqs. without polluting groundwater by mixing them with inventive absorbent blend) ΙT 1313-82-2, Sodium sulfide, reactions RL: RCT (Reactant); RACT (Reactant or reagent) (for complexation of heavy metals; disposal of solvent-containing ligs. without polluting groundwater by mixing them with inventive absorbent blend) HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1 373 ANSWERS L1CAPLUS COPYRIGHT 2007 ACS on STN CC 72-2 (Electrochemistry) Section cross-reference(s): 35, 36, 56 ΤI Electropolymerization of pyrrole on zinc-lead-silver alloys electrodes in acidic and neutral organic media ST electropolymn pyrrole zinc lead silver alloy electrode org media; sulfide pretreated zinc alloy electrode electropolymn pyrrole IT X-ray diffraction (by zinc-lead-silver alloys in pyrrole electropolymn. study) IT Polymerization (electrochem.; of pyrrole on zinc-lead-silver alloy electrodes in acidic and neutral organic media containing p-toluenesulfonate and effect of sulfide pretreatment of electrode) IT Cyclic voltammetry (in pyrrole electropolymn. on sulfide-pretreated zinc-lead-silver alloy in acetonitrile containing tetraethylammonium tosylate and pyrrole) IT Binding energy X-ray photoelectron spectra (of polypyrrole electrodeposited on pretreated zinc-lead-silver alloy electrodes) ITIR spectra (of polypyrrole films doped and undoped with p-toluenesulfonate) IT Adhesion, physical (of polypyrrole films electrosynthesized on pretreated zinc-lead-silver alloy) IT Polymer morphology (of polypyrrole films electrosynthesized on pretreated zinc-lead-silver alloy in organic media) IT Solvent effect (on electropolymn. of pyrrole on zinc-lead-silver alloy electrodes) IT Electric potential (oscillations; in pyrrole electropolymn. on zinc-lead-silver alloy in acetonitrile containing tetraethylammonium tosylate and pyrrole) IT Electron donors (solvent; in electropolymn. of pyrrole on zinc-lead-silver alloy electrodes in organic media containing p-toluenesulfonate) IT Chemically modified electrodes (sulfide-pretreated zinc alloy for pyrrole electropolymn.) IT 30604-81-0P, Polypyrrole RL: PNU (Preparation, unclassified); PRP (Properties); PREP (Preparation) (electrochem. preparation on zinc-lead-silver alloy electrodes and XPS spectra of polypyrrole film electrodeposited on pretreated zinc alloy)

```
477946-11-5, Lead 10, silver 25, zinc 65 477946-12-6, Lead 60, silver
IT
     10, zinc 30
     RL: DEV (Device component use); PRP (Properties); USES (Uses)
        (electrode with and without sulfide pretreatment for electropolymn. of
        pyrrole in acidic and neutral organic media containing p-toluenesulfonate)
IT
     7440-06-4, Platinum, uses
     RL: DEV (Device component use); PRP (Properties); USES (Uses)
        (electropolymn. of pyrrole on zinc-lead-silver alloy electrodes and
        comparison with platinum electrodes in organic media containing
        p-toluenesulfonate)
     75-05-8, Acetonitrile, uses
ΙT
     RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses)
        (electropolymn. of pyrrole on zinc-lead-silver alloy electrodes in
        acetonitrile containing tetraethylammonium p-toluenesulfonate and effect of
        sulfide pretreatment of electrode)
IT
     109-97-7, Pyrrole
     RL: PRP (Properties); RCT (Reactant); RACT (Reactant or reagent)
        (electropolymn. of pyrrole on zinc-lead-silver alloy electrodes in
        acidic and neutral organic media containing p-toluenesulfonate and effect of
        sulfide pretreatment of electrodes)
IT
     733-44-8, Tetraethylammonium p-toluenesulfonate
     RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses)
        (electropolymn. of pyrrole on zinc-lead-silver alloy electrodes in
        acidic and neutral organic media containing tetraethylammonium
        p-toluenesulfonate and effect of sulfide pretreatment of electrode)
IT
     98-95-3, Nitrobenzene, uses
     RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses)
        (electropolymn. of pyrrole on zinc-lead-silver alloy electrodes in
        nitrobenzene containing tetraethylammonium p-toluenesulfonate and effect of
        sulfide pretreatment of electrode)
IT
     108-32-7, Propylene carbonate
     RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses)
        (electropolymn. of pyrrole on zinc-lead-silver alloy electrodes in
        propylene carbonate containing tetraethylammonium p-toluenesulfonate and
        effect of sulfide pretreatment of electrode)
IT
     16722-51-3, p-Toluenesulfonic acid, ion(1-), properties
     RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP
     (Physical process); PROC (Process)
        (polypyrrole doped with; IR spectra of electrogenerated)
IT
     1313-82-2, Sodium sulfide na2s, uses
     RL: MOA (Modifier or additive use); PRP (Properties); USES (Uses)
        (pretreatment of zinc-lead-silver alloy electrodes by aqueous
        sodium sulfide for electropolymn. of pyrrole on
        zinc-lead-silver alloy electrodes in acidic and neutral organic
        media containing p-toluenesulfonate)
```

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

- L1 373 ANSWERS CAPLUS COPYRIGHT 2007 ACS on STN
 CC 79-6 (Inorganic Analytical Chemistry)
 Section cross-reference(s): 59, 60, 61
 TI Determination of zinc in environmental samples by anodic stripping voltammetry
- ST zinc detn environmental sample anodic stripping voltammetry
 IT Air analysis
 Anodic stripping voltammetry

Environmental analysis

```
Oyster
     Wastewater treatment sludge
        (zinc determination in environmental samples by anodic stripping voltammetry
        with sulfide addition for removal of copper(II) ion interference)
IT
     7732-18-5, Water, analysis
     RL: AMX (Analytical matrix); ANST (Analytical study)
        (zinc determination in environmental samples by anodic stripping voltammetry
        with sulfide addition for removal of copper(II) ion interference)
     7440-66-6, Zinc, analysis
IT
     RL: ANT (Analyte); ANST (Analytical study)
        (zinc determination in environmental samples by anodic stripping voltammetry
        with sulfide addition for removal of copper(II) ion interference)
IT
     1313-82-2, Sodium sulfide, analysis
     RL: ARU (Analytical role, unclassified); ANST (Analytical study)
        (zinc determination in environmental samples by anodic stripping
        voltammetry with sulfide addition for removal of copper(II) ion
        interference)
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1
L1
      373 ANSWERS
                    CAPLUS COPYRIGHT 2007 ACS on STN
IC
     ICM H01M006-06
     ICS H01M004-42
     52-2 (Electrochemical, Radiational, and Thermal Energy Technology)
CC
TI
     Manufacture of zinc alkali battery
ST
     zinc anode battery potassium sulfide; surfactant gelled zinc battery anode
IT
        (battery, zinc, gelled, manufacture of, alkali metal sulfides and
        surfactants in)
IT
     135832-86-9
                   141709-68-4
                                 141709-69-5
                                                141709-70-8
                                                              141709-71-9
     RL: USES (Uses)
        (anodes, manufacture of gelled, alkali metal sulfides and surfactants in,
        for alkaline batteries)
IT
     1312-73-8, Potassium sulfide
                                    1313-82-2, Sodium sulfide
                    1344-08-7, Sodium sulfide (\overline{Na2(Sx)})
     (Na2S), uses
     12136-58-2, Lithium sulfide 37199-66-9, Potassium sulfide (K2(Sx))
     141656-79-3
     RL: USES (Uses)
        (in gelled zinc anode manufacture, for batteries)
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1
L1
      373 ANSWERS
                    CAPLUS COPYRIGHT 2007 ACS on STN
CC
     66-3 (Surface Chemistry and Colloids)
TΙ
     Surface analysis of polymer systems. 3. Zinc sulfide cadmium
     sulfide/Nafion
ST
     Nafion cadmium zinc sulfide surface structure
IT
     Surface structure
        (of Nafion films containing cadmium sulfide and zinc sulfide ppts., light
        effect on)
     Light, chemical and physical effects
        (on surface structure of Nafion films containing precipitated cadmium
sulfide and
        zinc sulfide in presence of aqueous sodium
        sulfide)
IT
     131\overline{3-82-2}, properties
     RL: PRP (Properties)
```

```
(irradiation of Nafion films with visible light in presence of aqueous,
surface
        structure in relation to)
     7783-06-4, uses and miscellaneous
IT
     RL: USES (Uses)
        (precipitation by, of cadmium sulfide and zinc sulfide in Nation films,
surface
        structure in relation to)
IT
     1314-98-3, properties
     RL: PRP (Properties)
        (precipitation of cadmium sulfide and, in Nafion films, surface structure in
        relation to)
IT
     1306-23-6, properties
     RL: PRP (Properties)
        (precipitation of zinc sulfide and, in Nafion films, surface structure in
        relation to)
     66796-30-3
IT
     RL: PRP (Properties)
        (surface structure of, containing precipitated cadmium sulfide and zinc
sulfide)
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1
      373 ANSWERS CAPLUS COPYRIGHT 2007 ACS on STN
L1
CC
     9 (Electric and Magnetic Phenomena)
TI
     Luminescence of low-temperature manganese-activated zinc sulfide phosphors
IT
        (impact, of ZnS containing Cu and Mn suspended in polystyrene)
IT
     Luminescence
     Phosphorescence
        (of zinc sulfide, containing Mn, and effect of preparation conditions
thereon)
TΥ
     7647-01-0P, Hydrochloric acid
     RL: PREP (Preparation)
        (from vinyl chloride polymers, zinc sulfide Mn-containing phosphor prepared
        in presence of, luminescence of)
     7439-96-5P, Manganese
IT
     RL: PREP (Preparation)
        (phosphors containing, luminescence and phosphorescence of, effect of
        preparation on)
IT
     1314-98-3P, Zinc sulfide
     RL: PREP (Preparation)
        (phosphors, includes scintillators, luminescence and phosphorescence of
        Mg-containing, and effect of preparation conditions thereon)
IT
     9003-53-6, Styrene polymers
        (zinc sulfide Cu- and Mn-doped phosphor suspended in, impact
        luminescence of)
     1313-82-2P, Sodium sulfide, Na2S
IT
                                         7722-84-1P,
     Hydrogen peroxide
     RL: PREP (Preparation)
        (zinc sulfide Mn-containing phosphor prepared in presence of,
        luminescence of)
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1
```

CAPLUS COPYRIGHT 2007 ACS on STN

L1

IC

373 ANSWERS

ICM C08L033-26

ICS C08F002-44; C08F002-02; C08F120-56; C08K009-04; G02B001-04

CC 37-6 (Plastics Manufacture and Processing) Section cross-reference(s): 73

TI Method for preparation of transparent polymer optical material having high nanoparticle content

ST sulfur nanoparticle acrylic polymer composite optical material prepn

IT Nanocomposites

Optical materials

Transparent materials

(preparation of transparent polymer optical material having high nanoparticle content)

IT 26793-34-0P, Dimethyl acrylamide homopolymer 29760-26-7P, Dimethyl acrylamide-styrene copolymer 31018-13-0P, N,N-Dimethylacrylamide-methyl methacrylate copolymer 132535-35-4P, N,N-Dimethylacrylamide-divinylbenzene-styrene copolymer 896109-99-2P, N,N-Dimethylacrylamide-triethylene glycol diacrylate copolymer

RL: IMF (Industrial manufacture); POF (Polymer in formulation); TEM

RL: IMF (Industrial manufacture); POF (Polymer in formulation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (preparation of transparent polymer optical material having high nanoparticle content)

IT 62-55-5, Thioacetamide 62-56-6, Thiourea, uses 106-45-6, p-Methyl thiophenol 557-34-6, Zinc acetate 628-52-4, Chromium diacetate 1313-82-2, Sodium sulfide, uses 4946-13-8, p-Ethyl thiophenol 7646-85-7, Zinc chloride, uses 7733-02-0, Zinc sulfate 7783-06-4, Hydrogen sulfide, uses 10049-05-5, Chromium dichloride 13637-61-1, Zinc perchlorate 13931-95-8, Chromium diperchlorate 30499-56-0, Mercaptomethylstyrene 53339-53-0 242791-24-8

RL: MOA (Modifier or additive use); USES (Uses) (preparation of transparent polymer optical material having high nanoparticle content)

IT 67-68-5, Dimethyl sulfoxide, uses 68-12-2, N,N-Dimethylformamide, uses 127-19-5, Dimethyl acetamide

RL: NUU (Other use, unclassified); USES (Uses)
(preparation of transparent polymer optical material having high
nanoparticle content)

IT 60-24-2, Mercaptoethanol 96-27-5, Thioglycerol 108-98-5, Thiophenol, uses 63947-56-8

RL: MOA (Modifier or additive use); USES (Uses) (surface modifier; preparation of transparent polymer optical material having high nanoparticle content)

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L1 373 ANSWERS CAPLUS COPYRIGHT 2007 ACS on STN

CC 72-8 (Electrochemistry)
Section cross-reference(s): 56, 76, 78

TI Prospective trends of galvano plastics metallization of wax

ST galvanoplastics wax metalization nickel electrodeposition copper zinc sulfide deposition

IT Coating process

(electroless; of zinc and copper sulfides on wax surface in process of its metalization)

IT Coating process

(metalization; prospective trends of galvano plastics metalization of wax)

IT Electric conductivity
 (of copper and zinc sulfide film deposited on wax surface)

```
IT
    рН
        (of electrolyte for electrodeposition of nickel on sulfide activated
        surface of wax)
IT
     Electrodeposition
        (of nickel on sulfide activated surface of wax)
ΙT
    Waxes
     RL: MSC (Miscellaneous)
        (prospective trends of galvano plastics metalization of)
ΙT
     Electroforming
        (prospective trends of galvano plastics metalization of wax)
IT
     1336-21-6, Ammonium hydroxide 7733-02-0, Zinc sulfate
     Copper sulfate, reactions
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
    process); RCT (Reactant); PROC (Process); RACT (Reactant or reagent)
        (chemical deposition of zinc and copper sulfides on wax surface in process
        of its metalization, in solution containing)
IT
     1313-82-2, Sodium sulfide, reactions
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); RCT (Reactant); PROC (Process); RACT (Reactant or reagent)
        (chemical deposition of zinc and copper sulfides on wax surface
       in sulfate solution using)
IT
     7718-54-9, Nickel chloride, reactions 13770-89-3, Nickel Sulfamate
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); RCT (Reactant); PROC (Process); RACT (Reactant or reagent)
        (electrodeposition of nickel on sulfide activated surface of wax in
        solution containing)
     151-21-3, Sodium dodecylsulfate, uses
IT
                                             10043-35-3, Boric acid, uses
     RL: NUU (Other use, unclassified); USES (Uses)
        (electrodeposition of nickel on sulfide activated surface of wax in
        solution containing)
     7440-02-0P, Nickel, processes
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PNU (Preparation, unclassified); PREP (Preparation); PROC
     (Process)
        (prospective trends of galvano plastics metalization of wax.with)
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1
     . 373 ANSWERS
                    CAPLUS COPYRIGHT 2007 ACS on STN
L1
     72-8 (Electrochemistry)
     Section cross-reference(s): 55, 66
     Electroplating of plastics using a current-conducting layer
ST
     iron nickel alloy electrodeposition plastic current conducting layer
     Heat treatment
IT
        (of Fe-Ni alloy electrodeposits on plastics with current-conducting
        layer)
IT
     Adhesion, physical
     Mechanical properties
        (of Fe-Ni alloy electrodeposits with plastics with current-conducting
        layer)
IT
     Adsorption
        (of ammonia zinc complex on plastic surface during during plastics
        electroplating using current-conducting layer)
IT
     Current efficiency
        (of electrodeposition of Fe-Ni alloy on plastics with
        current-conducting layer from sulfamate solution)
IT
     Optimization
```

(of etching of plastics surface with CrO3-H2SO4 solution in process of electroplating layer)

IT Etching

(of plastics surface with CrO3-H2SO4 solution in process of electroplating layer)

IT Sulfidation

(of zinc hydroxide with Na2S on plastic during its electroplating using current-conducting layer)

IT Exchange reaction

(of zinc sulfide with Pb(NO3)2 during plastics electroplating using current-conducting layer)

IT Contact angle

(optimization of etching of plastics surface with CrO3-H2SO4 solution in process of electroplating layer, using)

RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses) (electrodeposition of Fe-Ni alloy on plastics with current-conducting layer from solution containing)

IT 7718-54-9, Nickel dichloride, properties 13770-89-3, Nickel sulfamate 14017-39-1, Iron sulfamate

RL: PRP (Properties); RCT (Reactant); RACT (Reactant or reagent) (electrodeposition of Fe-Ni alloy on plastics with current-conducting layer from solution containing)

IT 11148-32-6

RL: FMU (Formation, unclassified); PRP (Properties); FORM (Formation, nonpreparative)

(electroplating of plastics using current-conducting layer)

IT 9003-18-3, Acrylonitrile butadiene copolymer

RL: PRP (Properties)

(electroplating using current-conducting layer)

IT 1333-82-0, Chromium trioxide 7664-93-9, Sulfuric acid, uses
RL: NUU (Other use, unclassified); PRP (Properties); USES (Uses)
 (etching of plastics surface with CrO3-H2SO4 solution in process of electroplating layer)

IT 10099-74-8, Lead nitrate

RL: PRP (Properties); RCT (Reactant); RACT (Reactant or reagent) (exchange reaction with zinc sulfide during plastics electroplating using current-conducting layer)

IT 1314-87-0, Lead sulfide

RL: FMU (Formation, unclassified); PRP (Properties); FORM (Formation, nonpreparative)

(formation by exchange reaction on plastics surface during its electroplating using current-conducting layer from)

IT 1314-98-3, Zinc sulfide, properties 20427-58-1, Zinc hydroxide
RL: FMU (Formation, unclassified); PRP (Properties); FORM (Formation, nonpreparative)

(formation on plastic during its electroplating using current-conducting layer)

IT 1313-82-2, Sodium sulfide, properties

RL: PRP (Properties); RCT (Reactant); RACT (Reactant or reagent) (sulfurization of zinc hydroxide with Na2S on plastic during its electroplating using current-conducting layer)

```
L1
      373 ANSWERS
                  CAPLUS COPYRIGHT 2007 ACS on STN
IC
     ICM C01B017-22
     ICS C01B017-36
CC
     49-5 (Industrial Inorganic Chemicals)
     Manufacture of purified sodium sulfide from anhydrous mirabilite
ΤΊ
ST
     sodium sulfide manuf anhyd mirabilite; redn mirabilite coal sodium sulfide
     manuf
IT
     Coal
     RL: USES (Uses)
        (reductant, in sodium sulfide manufacture from anhydrous mirabilite)
IT
     1344-28-1P, Alumina, uses
     RL: PREP (Preparation); USES (Uses)
       (activated, iron removal by, in sodium sulfide manufacture from anhydrous
     14567-58-9, Mirabilite
IT
     RL: USES (Uses)
        (anhydrous, reduction of, with coal, in sodium sulfide manufacture)
     1305-78-8, Calcia, uses 1313-13-9, Manganese dioxide, uses
     Zinc oxide, uses 7601-54-9, Trisodium phosphate
     RL: USES (Uses)
        (iron removal by, in sodium sulfide manufacture from
        anhydrous mirabilite)
     1313-82-2P, Sodium sulfide, preparation
IT
     RL: IMF (Industrial manufacture); PREP (Preparation)
        (manufacture of, from mirabilite, by reduction using coal)
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1
      373 ANSWERS
                    CAPLUS COPYRIGHT 2007 ACS on STN
CC
     54-1 (Extractive Metallurgy)
ΤI
     Effect of oxygen on the behavior of reagents in the desorption of the
ST
     copper zinc flotation collector desorption
IT
     Zinc ores, preparation
     RL: PEP (Physical, engineering or chemical process); PROC (Process)
        (copper-, flotation of, xanthate desorption by sodium
        sulfide and activated carbon in)
     Copper ores, preparation
IT
     RL: PEP (Physical, engineering or chemical process); PROC (Process)
        (zinc-, flotation of, xanthate desorption by sodium
        sulfide and activated carbon in).
     7440-44-0, Carbon, uses and miscellaneous
IT
     RL: USES (Uses)
        (activated, desorption by, of xanthate collector in copper-zinc ore
        flotation)
IT
     1313-82-2, Sodium sulfide, uses and miscellaneous.
     RL: PEP (Physical, engineering or chemical process); PROC (Process)
        (desorption with, of xanthate collector in copper-zinc ore
        flotation)
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0
=> zinc (S) remov? (S) "sodium sulfide"
       .641794 ZINC
           139 ZINCS
        641819 ZINC
                 (ZINC OR ZINCS)
```

```
1317840 REMOV?
       1147007 "SODIUM"
            40 "SODIUMS"
       1147016 "SODIUM"
                 ("SODIUM" OR "SODIUMS")
        332880 "SULFIDE"
         87975 "SULFIDES"
        366071 "SULFIDE"
                 ("SULFIDE" OR "SULFIDES")
         13224 "SODIUM SULFIDE"
                 ("SODIUM"(W) "SULFIDE")
L2
            30 ZINC (S) REMOV? (S) "SODIUM SULFIDE"
=> d scan
L2
      30 ANSWERS
                   CAPLUS COPYRIGHT 2007 ACS on STN
     C01F007-02A; C01F007-34B; C01G003-12B
IC
INCL 023141000
     54 (Extractive Metallurgy)
     Purifying aluminate solutions
ST
     aluminate soln purifn; refining aluminate soln; zinc removal aluminate
IT
     7440-66-6, uses and miscellaneous
     RL: REM (Removal or disposal); PROC (Process)
        (removal of, from aluminate solns. by sodium sulfide)
IT
     1344-28-1P, preparation
     RL: PREP (Preparation)
        (zinc removal from aluminate solns. in, by
        sodium sulfide)
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1
L2
      30 ANSWERS
                   CAPLUS COPYRIGHT 2007 ACS on STN
CC
     63-4 (Pharmaceuticals)
     Method for producing active polypeptide-calcium and zinc-protein from sea
ST
     polypeptide calcium zinc protein sea cucumber
IT
     Centrifugation
     Freezing
     Holothurioidea
     Homogenization
     Polymerization
     Pulping
        (method for producing active polypeptide-calcium and zinc-protein from
        sea cucumber)
IT
     Peptides, biological studies
     RL: FFD (Food or feed use); IMF (Industrial manufacture); RCT (Reactant);
     BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent);
     USES (Uses)
        (method for producing active polypeptide-calcium and zinc-protein from
        sea cucumber)
ΙT
     Proteins
     RL: FFD (Food or feed use); RCT (Reactant); BIOL (Biological study); RACT
     (Reactant or reagent); USES (Uses)
        (method for producing active polypeptide-calcium and zinc-protein from
        sea cucumber)
     9002-07-7, Trypsin
ΙT
```

```
RL: CAT (Catalyst use); USES (Uses)
         (method for producing active polypeptide-calcium and zinc-protein from
        sea cucumber)
     7440-66-6, Zinc, biological studies 7440-70-2, Calcium, biological
IT
     RL: FFD (Food or feed use); RCT (Reactant); BIOL (Biological study); RACT
      (Reactant or reagent); USES (Uses)
         (method for producing active polypeptide-calcium and zinc-protein from
        sea cucumber)
     64-17-5, Ethanol, uses 64-19-7, Acetic acid, uses
IT
                                                            1313-82-2. Sodium
     sulfide, uses
     RL: NUU (Other use, unclassified); USES (Uses)
         (method for producing active polypeptide-calcium and zinc-protein from
        sea cucumber)
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1
                   CAPLUS COPYRIGHT 2007 ACS on STN
L2
      30 ANSWERS
CC.
     54-2 (Extractive Metallurgy)
TI.
     Electrolytic method for preparing zinc by alkaline leaching process
     electrolysis zinc alk leaching calcium oxide sodium sulfide
ST
IT
     Glues
         (bone glues; electrolytic method for preparing zinc by alkaline leaching
        process)
ΙT
     Smelting
         (electrolytic method for preparing zinc by alkaline leaching process)
IT
         (glues; electrolytic method for preparing zinc by alkaline leaching process)
IT
     7440-44-0, Carbon, uses
     RL: NUU (Other use, unclassified); USES (Uses)
         (activated; electrolytic method for preparing zinc by alkaline leaching
        process)
     1314-13-2, Zinc oxide, processes 3486-35-9, Zinc carbonate 14374-77-7,
IT
     Willemite
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
         (electrolytic method for preparing zinc by alkaline leaching process)
 IΤ
     7439-92-1, Lead, processes
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); REM (Removal or disposal); PROC (Process)
         (electrolytic method for preparing zinc by alkaline leaching process)
     1305-78-8, Calcium oxide, uses 1310-73-2, Sodium hydroxide, uses
 IT
      1313-82-2, Sodium sulfide, uses 1663-45-2, DPE
                                                         902759-57-3, JZ 1
      RL: NUU (Other use, unclassified); USES (Uses)
         (electrolytic method for preparing zinc by alkaline leaching process)
 IT
      7440-66-6P, Zinc, preparation
      RL: SPN (Synthetic preparation); PREP (Preparation)
         (electrolytic method for preparing zinc by alkaline leaching process)
      12597-68-1, Stainless steel, uses
· IT
      RL: TEM (Technical or engineered material use); USES (Uses)
         (electrolytic method for preparing zinc by alkaline leaching process)
```

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

- L2 30 ANSWERS CAPLUS COPYRIGHT 2007 ACS on STN
- CC 54-2 (Extractive Metallurgy)
- TI Combined wet method-dry method for recovering copper, nickel, and noble

metals from heavy metal-containing sludge resulted from production of electronics and electroplating

ST recovery copper nickel noble metal waste

IT Extraction

Leaching

Precipitation (chemical)

Sintering Smelting

Wastes

(combined wet method-dry method for recovering copper, nickel, and noble metals from heavy metal-containing sludge resulted from production of electronics and electroplating)

TT 7440-02-0P, Nickel, preparation 7440-05-3P, Palladium, preparation 7440-22-4P, Silver, preparation 7440-50-8P, Copper, preparation

7440-57-5P, Gold, preparation

RL: PUR (Purification or recovery); PREP (Preparation)
(combined wet method-dry method for recovering copper, nickel, and
noble metals from heavy metal-containing sludge resulted from production of
electronics and electroplating)

IT 497-19-8, Sodium carbonate, reactions 1313-82-2, Sodium sulfide,
 reactions 7664-93-9, Sulfuric acid, reactions 7681-49-4, Sodium
 fluoride, reactions 7722-84-1, Hydrogen peroxide, reactions
 RL: RGT (Reagent); RACT (Reactant or reagent)
 (combined wet method-dry method for recovering copper, nickel, and
 noble metals from heavy metal-containing sludge resulted from production of

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

electronics and electroplating)

=> d his

(FILE 'HOME' ENTERED AT 15:37:39 ON 11 OCT 2007)

FILE 'CAPLUS' ENTERED AT 15:37:51 ON 11 OCT 2007

L1 373 ZINC (S) "SODIUM SULFIDE"

L2 30 ZINC (S) REMOV? (S) "SODIUM SULFIDE"

=> d 12 1-30 ibib abs

L2 ANSWER 1 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2007:467752 CAPLUS

DOCUMENT NUMBER:

146:507539

TITLE:

Method for producing active polypeptide-calcium and

zinc-protein from sea cucumber

INVENTOR(S):

Ji, Jianguo; Li, Guozhong; Huang, Shikuan

PATENT ASSIGNEE(S):

Peop. Rep. China

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 8pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent Chinese

LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	_			
CN 1951402	Α	20070425	CN 2006-10069408	20061013
PRIORITY APPLN. INFO.:			CN 2006-10069408	20061013

The title method comprises: (1) removing intestine of fresh sea AΒ cucumber, washing, pulping, adding distilled water at a pulp/water volume ratio of 1:1, adjusting pH to 4.5-5.5 with glacial acetic acid, and stirring for 1-3 h, (2) adding sodium sulfide to pH 5-6, filtering, adding trypsin (15-25 weight% of the fresh sea cucumber), and homogenizing for reaction at 40-45°C for 6-10 h, (3) adding biol. calcium (3-10 weight% of the fresh sea cucumber) and/or biol. zinc (0.5-1 weight% of the fresh sea cucumber), and polymerizing to form polypeptide-calcium and/or zinc-protein, and (4) adding 3 times of ethanol, freezing at -5 to -15°C for 8-10 h, centrifuging, and vacuum-drying at 5-40°C to obtain the product. The invention has the advantages of calcium supplement, skeleton structure improvement, radiation and UV irradiation resistance, intravital metallic toxin removal, proper nutrient combination, and simple process.

ANSWER 2 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2006:753796 CAPLUS

DOCUMENT NUMBER:

145:192390

TITLE:

Electrolytic method for preparing zinc by alkaline

leaching process

INVENTOR(S):

Kuang, Zhengguo

PATENT ASSIGNEE(S):

Peop. Rep. China

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 7 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent

LANGUAGE:

Chinese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1807665	A .	20060726	CN 2005-10010613	20050118
PRIORITY APPLN. INFO.:			CN 2005-10010613	20050118

AB The title method comprises: (1) pulverizing zinc raw ore, adding a separating agent (CaO) 30 weight% of the dry total ore, milling, pretreating in.

a pretreating kettle at 40-50° for 50 min, and filtering to remove the pretreating solution, (2) feeding into a leaching kettle, adding sodium hydroxide (the ratio of sodium hydroxide to zinc = 6:1), leaching at $75-85^{\circ}$ for 35-40 min, and filtering, (3)

purifying the filtrate at normal temperature for 30-35 min by adding separating agent

(sodium sulfide) 4 times of the Pb content in the filtrate, and filtering, and (4) feeding the filtrate into an electrolyte tank, feeding into an electrolytic bath, adding additives such as bone glue 0.5-0.6 kg per ton zinc, regulating alkaline/zinc ratio to 7-10, and electrolyzing. The method can use lean ore such as zinc oxide ore, zinc carbonate ore, galmey, willemite, etc. with zinc contents less than 20%, and can reduce the cost and alkaline consumption.

ANSWER 3 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2006:429611 CAPLUS

DOCUMENT NUMBER:

145:149507

TITLE:

Combined wet method-dry method for recovering copper, nickel, and noble metals from heavy metal-containing sludge resulted from production of electronics and

electroplating

INVENTOR(S):

Sun, Tao; Yue, Xilong

PATENT ASSIGNEE(S):

Peop. Rep. China

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 17 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent

LANGUAGE:

Chinese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1718784	Α	20060111	CN 2005-10035633	20050707
PRIORITY APPLN. INFO.:		•	CN 2005-10035633	20050707

The title process includes leaching sludge with sulfuric acid in a reactor; adding sodium sulfide into the reactor to form copper sulfide entering into copper-containing sludge; adding iron powder to replace copper, noble metals gold, silver, palladium entering into copper-containing sludge; filtering to give nickel-containing solution and copper-containing sludge; oxidizing bivalent iron ions in the nickel solution into trivalent iron ions with iron oxidizer, neutralizing with sodium carbonate to precipitate iron, filtering to remove iron as well as chromium; adding sodium fluoride to remove impurities calcium, magnesium, and aluminum etc., wherein iron and less than 5% nickel enter nickel-containing sludge; further extracting iron, copper, chromium, calcium, magnesium, aluminum, zinc from remaining nickel solution, reverse extracting resulted organic phase using sulfuric acid to get pure nickel solution;

precipitating nickel using sodium carbonate to get refined nickel carbonate; resp.

sintering the copper-containing sludge and the nickel-containing iron sludge to get sintered copper cake and sintered nickel cake; and reducing/smelting the copper cake and the nickel cake at high temperature to obtain black copper and low-grade nickel matte, wherein the copper and noble metals gold, silver, palladium are recovered from copper black, and the nickel is recovered from low-grade nickel matte.

ANSWER 4 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2005:693118 CAPLUS

DOCUMENT NUMBER:

144:356896

TITLE:

Gamma ray-induced reduction and removal of heavy metal

ions in aqueous solutions

AUTHOR(S):

Wu, Xing-Zheng; Takami, Takayuki; Somekawa, Kenichi;

Nishio, Shigeru

CORPORATE SOURCE:

Department of Materials Science and Engineering,

Faculty of Engineering, Fukui University, Fukui-shi,

910-8507, Japan

SOURCE:

Radioisotopes (2005), 54(6), 179-184

CODEN: RAISAB; ISSN: 0033-8303

PUBLISHER:

Nippon Aisotopu Kyokai

DOCUMENT TYPE:

Journal

English

LANGUAGE: Possible reduction and removal of heavy metals (Ag+, Zn2+, Cu2+, Co2+) from

water by γ -ray irradiation was examined No concentration change was observed when aqueous solns. of metal ions were irradiated with γ -rays directly or

after bubbling with He gas. Metal concns. were reduced to lower oxidation states by irradiation in the presence of Na2SO3. Ppts. were observed in all irradiated solns. containing 4 metals (Ag+, Zn2+, Cu2+, Co2+). Ag+ was reduced to metallic Ag.

REFERENCE COUNT:

12

THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L2 ANSWER 5 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:258983 CAPLUS

DOCUMENT NUMBER:

141:263684

TITLE:

Flotation study on separating skarn-type complicated

copper-zinc sulphide ore Yu, Xue; Ma, Guang-qing

AUTHOR(S):
CORPORATE SOURCE:

Shenyang Research Institute of Nonferrous Metals,

Shenyang, 110023, Peop. Rep. China

SOURCE:

Youse Jinshu, Xuankuang Bufen (2003), (6), 10-13

CODEN: YJXBA5; ISSN: 1671-9492

PUBLISHER:

Youse Jinshu Bianjibu

DOCUMENT TYPE:

Journal

LANGUAGE:

Chinese

AB The view of using sodium sulfide to remove

and fix Cu2+ in the pulp (sodium sulfide has function

of restrainer of sphalerite and pyrite and removing reagent),

using SK9011 that has more strong selecting as collectors of separating copper

and $\underline{\mathbf{zinc}}$, using reasonable proportion of $\underline{\mathbf{zinc}}$ sulfate

and sodium sulfite as main restrainer of sphalerite was advanced, adopting prior-pulp flotation technol. flowsheet aiming at characters of the

Complicated skarn type copper-zinc sulfide ore of closing

dissemination size and sphalerite being activated Cu2+, the separating copperzinc question of this ore was solved.

L2 ANSWER 6 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2002:669405 CAPLUS

DOCUMENT NUMBER:

137:190298

TITLE:

Removal of chromium VI compounds and

zinc complexes from metalworking wastewater by

using waste acid and waste sodium

sulfide

INVENTOR(S):

Gosch, Bodo

PATENT ASSIGNEE(S):

Sachtleben Chemie GmbH, Germany

SOURCE:

Ger. Offen., 2 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

Campan

FAMILY ACC. NUM. COUNT:

German

PAMILI ACC. NOM. COON

1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	DE 10063580	A1	20020905	DE 2000-10063580	20001220
PRIOR	RITY APPLN. INFO.:			DE 2000-10063580	20001220

AB Cr VI compds. and Zn chelate complexes are removed from metalworking wastewater by treatment with waste acid containing H2SO4 and FeSO4 followed by treatment with waste Na2S and neutralization with NaOH or lime milk.

REFERENCE COUNT:

1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L2 ANSWER 7 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2001:777548 CAPLUS

Correction of: 2000:803419

DOCUMENT NUMBER:

135:291721

Correction of: 134:59355

TITLE:

Integrated hydrometallurgical process for production of zinc from electric arc furnace dust in alkaline

medium

AUTHOR(S):

Zhao, Youcai; Stanforth, Robert

CORPORATE SOURCE:

Department of Environmental Engineering, National University of Singapore, Singapore, 119260, Singapore

SOURCE:

Journal of Hazardous Materials (2000), 80(1-3),

223-240

CODEN: JHMAD9; ISSN: 0304-3894

PUBLISHER:

Elsevier Science B.V.

DOCUMENT TYPE:

Journal

English LANGUAGE:

AB. In this study, a novel and integrated hydrometallurgical process for the production of zinc powder from elec. arc furnace (EAF) dust in alkaline medium is

reported. The dust is firstly hydrolyzed in water, and then fused in caustic soda at 350° for 1 h, followed by leaching in alkaline solution in which both zinc and lead are effectively extracted Zinc powder is then produced by electrowinning from the leach solution after the lead is selectively removed by precipitation using sodium sulfide as precipitant. The EAF dust tested contained 25% Zn, 1.8% Pb and 33% Fe. It was found that 38% of zinc and 68% of lead could be extracted from the dust when leached directly in caustic soda solution Leaching of zinc increased to 80% when dust was directly fused with caustic soda followed by alkaline leaching. However, the leaching further increased to 95% when the dust was hydrolyzed first with water before fusion. Zinc powder with a purity of 99.95% was then produced by electrowinning from the lead depleted solution Stainless electrodes were

REFERENCE COUNT:

15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 8 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

used as both anode and cathode.

ACCESSION NUMBER:

2001:128679 CAPLUS

DOCUMENT NUMBER:

134:270557

TITLE:

Removal of zinc from rubber thread manufacturing

industry wastewater using chemical

precipitant/flocculant

AUTHOR(S):

Subbiah, R. M.; Sastry, C. A.; Agamuthu, P.

CORPORATE SOURCE:

Green and Smart Sdn. Bhd., Bangunan Mardec, Selangor

Darul Ehsan, 6800, Malay.

SOURCE:

Environmental Progress (2000), 19(4), 299-304

CODEN: ENVPDI; ISSN: 0278-4491

PUBLISHER:

American Institute of Chemical Engineers

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Wastewater from the rubber thread manufacturing industry is acidic in nature with pH 3.8-4.4 and containing high concns. of COD, BOD, total-N and Zn. The high BOD: COD ratio of the wastewater (0.78-0.90) indicates that the rubber wastewater is biodegradable and can be treated effectively by an anaerobic digestion process commonly used in this industry. However, because the wastewater contained high concns. of Zn, 250-310 mg/L, which could inhibit the digestion process, it was pretreated by chemical

precipitation/flocculation. A

treatment process using a combination of Na sulfide and polyelectrolyte LT 27 at concns. 800 and 5 mg/L, resp., is the most cost-effective system for 2n removal from the wastewater. 2n removal by Na sulfide is not affected by pH. The optimum settling time was 60 min, whereas the optimum flocculation time was 20 min, and a speed of 20 rpm in a 110-mm diameter reactor gave the best results. The estimated cost of 2n removal using this method was 2n 2n 2n0 method was 2n0 method was

REFERENCE COUNT:

THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L2 ANSWER 9 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2000:803419 CAPLUS

DOCUMENT NUMBER:

134:59355

TITLE:

Integrated hydrometallurgical process for production

of zinc from electric arc furnace dust in alkaline

medium

AUTHOR(S):

Youcai, Z.; Stanforth, R.

CORPORATE SOURCE:

Department of Environmental Engineering, National

University of Singapore, Singapore, 119260, Singapore

SOURCE:

Journal of Hazardous Materials (2000), 80(1-3),

223-240

CODEN: JHMAD9; ISSN: 0304-3894

PUBLISHER:

Elsevier Science B.V.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB In this study, a novel and integrated hydrometallurgical process for the production of zinc powder from elec. arc furnace (EAF) dust in alkaline medium is

reported. The dust is firstly hydrolyzed in water, and then fused in caustic soda at 350° for 1 h, followed by leaching in alkaline solution in which both zinc and lead are effectively extracted **Zinc** powder is then produced by electrowinning from the leach solution after the lead is selectively **removed** by precipitation using **sodium**

sulfide as precipitant. The EAF dust tested contained 25% Zn, 1.8% Pb and 33% Fe. It was found that 38% of zinc and 68% of lead could be extracted from the dust when leached directly in caustic soda solution Leaching of zinc increased to 80% when dust was directly fused with caustic soda followed by alkaline leaching. However, the leaching further increased to 95% when the dust was hydrolyzed first with water before fusion. Zinc powder with a purity of 99.95% was then produced by electrowinning from the lead depleted solution Stainless electrodes were used as both anode and cathode.

REFERENCE COUNT:

THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L2 ANSWER 10 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:734152 CAPLUS

DOCUMENT NUMBER:

131:312204

TITLE:

Manufacture of high-purity active zinc oxide using

circulating ammonia water for impurity removal

INVENTOR(S):

Li, Liben; Yang, Guohua; Dai, Shubin

PATENT ASSIGNEE(S):

Huaxin Zinc Products Factory, Jintan City, Peop. Rep.

China

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 10 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent

LANGUAGE:

Chinese

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1149552	` A	19970514	CN 1995-112746	19951027
PRIORITY APPLN. INFO.:			CN 1995-112746	19951027

AB The process comprises (1) leaching crude ZnO (ZnO content >60%) with NH4OH, and NH4HCO3 (ZnO: NH4OH: NH4HCO3 weight ratio 1:8:(1-1.5)) at 50-80°, (2) adjusting the pH to neutrality and filtering, (3) reducing the impurities in the solution in two step process with H2O2 and Zn dust and then forming sulfide by adding Na2S, (4) evaporating for 5-8 h at $65-95^{\circ}$ and from -0.05 to 0 MPa (ammonia obtained is absorbed by water and recycled), (5) dewatering, (6) drying at 110-150° for 2-3 h to obtain the basic Zn carbonate, and (7) calcining at 450-550°.

L2ANSWER 11 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:381141 CAPLUS

DOCUMENT NUMBER:

131:75669

TITLE:

Purification and impurity removing in production of

active zinc oxide by coordination

AUTHOR(S):

Peng, Xiaochun; Peng, Qingjing; Zou, Xiaoyong; Huang,

CORPORATE SOURCE:

Department of Chemistry, Jishou University, Jishou,

416000, Peop. Rep. China

SOURCE:

Wujiyan Gongye (1999), 31(3), 23-24

CODEN: WUGOFJ; ISSN: 1006-4990

PUBLISHER:

Wujiyan Gongye Bianjib

DOCUMENT TYPE:

Journal

LANGUAGE: Chinese

Active ZnO was prepared by coordination method by making processes of extracting

and impurity removing be carried out simultaneously in production process by using extracting solution containing NH3 and NH4+ and Na2S as impurity remover. Impurities of Pb, Ca, and Cu in raw material were converted into sulfides, which were not soluble in ammonia liquor, thus were separated from Zn by filtration. Effect of Na2S amount on impurity removing effect and effect of removing impurities by Na2S on leaching rate of Zn and total yield were discussed.

ANSWER 12 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:250 CAPLUS

DOCUMENT NUMBER:

130:143502

TITLE:

Study on treating the acidic waste at Northwest Lead

and Zinc Plant

AUTHOR(S):

Wang, Xinwen

CORPORATE SOURCE:

Northwest Copper and Lead Refining Plant, Baiyin,

730900, Peop. Rep. China

SOURCE:

Youse Jinshu (1998), 50(4), 126-129, 132

CODEN: YSCSAE; ISSN: 1001-0211

PUBLISHER:

Youse Jinshu Bianjibu

DOCUMENT TYPE:

Journal

LANGUAGE:

Chinese

A 2-stage neutralization (with slaked lime) flocculation (with S-3) process for treating acidic and smelting wastewaters showed effective heavy metal removal.

L2 ANSWER 13 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1998:704999 CAPLUS

DOCUMENT NUMBER: 130:60293

TITLE: Determination of zinc in environmental samples by

anodic stripping voltammetry

AUTHOR(S): Lau, Oi-Wah; Cheng, Oi-Ming

CORPORATE SOURCE: Department of Chemistry, The Chinese University of

Hong Kong, Shatin, Hong Kong

SOURCE: Analytica Chimica Acta (1998), 376(2), 197-207

CODEN: ACACAM; ISSN: 0003-2670

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

The copper(II) ion interference in zinc determination by anodic stripping voltammetry is removed by the addition of sulfide. The proposed method is applicable to a number of environmental samples. The calibration graph was linear from 0.3 to 11 μg of zinc. The recovery of zinc in three real samples and two synthetic sea water samples spiked with 10 trace metals using the proposed method ranged from 98% to 102%, and the precision of the method was 1.6% (n = 10) at the 1 μg level of zinc. Most of the metal ions and anions, except cobalt and nickel, found in the samples under study were found not to interfere with the determination. The method is simple, efficient and accurate, and was used for the determination of zinc in a number of natural water, domestic wastewater, air, oyster tissue, sewage sludge and sediment samples.

REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L2 ANSWER 14 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1997:793487 CAPLUS

DOCUMENT NUMBER: 128:37658

TITLE: Fundamental studies on the removal of copper, tin and

zinc from remelted steel scrap

AUTHOR(S): Nagasaka, T.; Hino, M.; Ban-Ya, S.

CORPORATE SOURCE: Department of Metallurgy, Faculty of Engineering,

Tohoku University, Sendai, 980-77, Japan

SOURCE: Process Technology Conference Proceedings (1997), 15,

41-49

CODEN: PTECEO; ISSN: 1075-8607

PUBLISHER: Iron and Steel Society

DOCUMENT TYPE: Journal LANGUAGE: English

AB Some laboratory studies have been done to discuss the possibility of copper, tin, and zinc removal from molten steel scrap. At first, the equilibrium copper distribution ratio between FeS-based sulfide flux and Fe-Csat melt at 1673K is reviewed. The copper distribution ratio, LCu, increases with the addition of alkaline or alkaline earth sulfide into FeS and reaches the

while the sulfur content in metal decreases at the same time. The maximum value of LCu of approx. 20-30 is obtained in FeS-M2S (M: Li, Na, and K) and FeS-MS (M: Sr and Ba) fluxes. In the second, the vaporization rate of zinc from liquid Fe-Zn alloy is discussed at 1873K by impinging Ar onto the surface of inductively stirred melt. The evaporation rate of zinc is found to be mainly controlled by the liquid phase mass transfer under the exptl. conditions of gas velocity >5 L/min. The evaporation rate of tin from Fe-S-CSat bath is also studied at 1673K under reduced pressure. The evaporation rate of tin is controlled mainly by the gas phase mass transfer and

becomes drastically faster by adding sulfur in the metal, and it indicates that tin dissolved in liquid iron could be removed by evaporation in the form of

SnS. The removal rate of tin, zinc, or other residual elements from molten steel in the conventional practical operations or vacuum treatment is also discussed based on the present results.

REFERENCE COUNT:

20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 15 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN L2

ACCESSION NUMBER:

1996:619880 CAPLUS

DOCUMENT NUMBER:

125:281220

TITLE:

Arsenic distribution in copper smelter and an

environmentally friendly way to remove it

AUTHOR(S):

Vircikova, Edita; Imris, Ivan

CORPORATE SOURCE:

Faculty Metallurgy, Technical University, Kosice, 040

00, Slovakia

SOURCE:

Proceedings of the Copper 95--Cobre 95 International Conference, Santiago, Nov. 26-29, 1995 (1995), Volume 2, 347-360. Editor(s): Casali, Aldo C. Canadian Institute of Mining, Metallurgy and Petroleum:

Montreal, Que. CODEN: 63KSAV

DOCUMENT TYPE:

Conference English

LANGUAGE:

AΒ The material balance of arsenic during pyrometallurgical production of copper in the Copper Smelter Krompachy, Slovakia, suggests that arsenic is an extremely harmful element for the copper-making process and it should be removed as much as possible from the concentrate before its smelting to matte, or from the converter flue dusts where most of the arsenic accumulates during the converting process. The converter flue dust is a serious problem in terms of pollution and storage because of the high content of metal in water-soluble compds. and because of its low apparent d. The pyrometallurgical treatment of the flue dusts gives no good yields because speiss is formed and it causes some troubles in the operations. This paper describes a laboratory-scale study of a hydrometallurgical method for processing of the flue dust from a converter in the Copper Smelter Krompachy (Slovakia) with the special aim to remove the arsenic. In order to overcome the difficulties, a hydrometallurgical treatment of converter flue dusts was developed at the Faculty of Metallurgy, Tech. University, Kosice. The process consists of a selective leaching of the flue dust in which arsenic is dissolved in sodium sulfide solution and then removed by precipitation methods using various agents: zinc sulfate electrolyte, copper sulfate, mixture of phosphoric acid with calcium hydroxide, calcium oxide and ferrous or ferric sulfate. In principle, arsenic can be removed in the form of a marketable product (copper arsenic for wood preservation) or as stable precipitate compds. suitable for safe

ANSWER 16 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1994:195240 CAPLUS

DOCUMENT NUMBER:

120:195240

landfill disposal under existing EPA regulations.

TITLE:

Manufacture of purified sodium sulfide from anhydrous

mirabilite

INVENTOR(S): PATENT ASSIGNEE(S): Wang, Yongshan; Hu, Yili; Ji, Yongping Tianshan Chemical Plant, Peop. Rep. China

SOURCE:

Faming Zhuanli Shenging Gongkai Shuomingshu, 7 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent Chinese

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1075299	Α	19930818	CN 1992-114398	19921206
CN 1033156	В	19961030		

PRIORITY APPLN. INFO.:

19921206 CN 1992-114398

AB The process comprises firing a mixture containing anhydrous mirabilite and coal,

impregnating the fired mixture, adding Fe-removing agent into the solution while stirring at 50-100°, separating the solution from solid, and cooling the solution to $20-45^{\circ}$ to obtain purified Na2S crystals. The products contain low Fe and other impurities such as Na2CO3 and NaCl.

ANSWER 17 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1993:479164 CAPLUS

DOCUMENT NUMBER:

119:79164

TITLE:

Calculation of some optimal pH values in wastewater

treatment by the use of equilibrium relations

AUTHOR(S):

Apak, Resat

CORPORATE SOURCE:

SOURCE:

Muhendislik Fak., Istanbul Univ., Istanbul, Turk. Kim. Kim. Muhendisligi Semp., 8th (1992), Volume 4, 349-53. Editor(s): Aydin, Adnan. Marmara Univ. Fac.

Sci. Lett.: Istanbul, Turk.

CODEN: 59AOAY

DOCUMENT TYPE:

. Conference

LANGUAGE:

Turkish

Optimal empirical pH values encountered in wastewater treatment processes, e.g., precipitation, flocculation and oxidation-reduction were determined in accord with

equilibrium calcns. The problems discussed are optimal pH in coaqulation processes carried out with Al and Fe(III) salts; pH of min. solubility of Al and Fe(III) phosphates in phosphate precipitation; optimal pH of Zn(II) treatment

by lime-Na precipitation; the pH of initiation of uranyl hydroxide precipitation as a

function of total uranyl concentration; Fe and Mn removal from groundwater by aeration followed by precipitation of hydrated oxides; optimal pH of total oxidation

of CN- by Cl.

ANSWER 18 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1993:171983 CAPLUS

DOCUMENT NUMBER:

118:171983

TITLE: INVENTOR(S): Manufacture of zinc oxide by the ammonia method Tang, Muotang; Lu, Junle; Yuan, Yansheng; et al. Zhongnan Polytechnic University, Peop. Rep. China

PATENT ASSIGNEE(S):

Faming Zhuanli Shenging Gongkai Shuomingshu, 7 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent Chinese

LANGUAGE:

SOURCE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1065685	Α	19921028	CN 1992-103230	19920428
CN 1028768	В	19950607		
PRIORITY APPLN. INFO.:			CN 1992-103230	19920428

AB Zn-containing raw materials are impregnated with a solution containing N 1.5-8.5 N,

NH4 salt 1.5-8.5 N, (NH4)2S2O8 0-10% of weight of raw material at a liquid/solid ratio of (2-10):1 and 0-90% for 0.5-4 h, and the impregnating solution is treated with S2- to remove other metal ions by forming metal sulfide. The purified Zn-containing solution is then treated with

a complex salt, e.g., ZnSO4.(NH4)2SO4, to produce Zn(OH)2 precipitate which is sintered at 300-900° for 0.25-6 h to produce high-purity ZnO.

L2 ANSWER 19 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1992:109539 CAPLUS

DOCUMENT NUMBER:

116:109539

TITLE:

Manufacture of zinc sulfide powder

INVENTOR(S):

Fujita, Kenji; Ogasawara, Tadashi; Natsume, Yoshitake

PATENT ASSIGNEE(S):

Osaka Titanium Co., Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 4 pp. CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 03232722	A	19911016	JP 1990-27951	19900207
PRIORITY APPLN. INFO.:			JP 1990-27951	19900207

AB In preparation of ZnS powder, aqueous solns. of Na2S and ZnCl2 are mixed to allow

reaction. Residual S-containing Na2S may be used for reaction with ZnCl2 to give ZnS suspension which is treated with NH3(g) to give ZnS powder. Preferably, the products are filtered, washed, and dried in H. The product is highly pure.

L2 ANSWER 20 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1990:577594 CAPLUS

DOCUMENT NUMBER:

113:177594

TITLE:

Sodium sulfide/ferrous sulfate metals treatment for

hazardous waste minimization

AUTHOR(S):

Carpenter, Charles; Suciu, Dan; Wikoff, Penny

CORPORATE SOURCE:

Air Force Eng. Serv. Cent., Tyndall Air Force Base,

FL, 32403, USA

SOURCE:

Proceedings of the Industrial Waste Conference (1990),

Volume Date 1989, 44th, 617-24 CODEN: PIWCAX; ISSN: 0073-7682

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB The Na2S/FeSO4 process is described for treatment of acid wastewaters from electroplating and metal finishing, which contain Cr, Ni, Cd, Cu, Pb, and Zn. The treatment involves neutralization of effluents with reduction of Cr(VI) and precipitation of the metals with sludge volume smaller than that obtained

by conventional neutralization/precipitation The Cr(VI) content in the effluent

is reduced to <0.1 mg/L at pH 7.2-7.5. The design and development of pilot- and full-scale installations, selection of operation parameters, and economic and environmental issues are described and discussed.

ANSWER 21 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1988:226315 CAPLUS

DOCUMENT NUMBER:

108:226315

TITLE:

An air stripping method for treatment of

electroplating solutions

AUTHOR(S):

Pearson, Donald E.; Bowers, Alan R.

CORPORATE SOURCE: SOURCE:

Viking Industries, Inc., Murfreesboro, TN, 37133, USA Hazardous Waste & Hazardous Materials (1988), 5(1),

85-91

CODEN: HWHME2; ISSN: 0882-5696

DOCUMENT TYPE:

Journal

LANGUAGE: English

The hypochlorite method of CN- removal from electroplating wastewaters (both real and simulated) is compared with that of air stripping for CNremoval and recovery as NaCN. The CN-, as HCN, is purged with air at .apprx.80 $^{\circ}$ and then absorbed in a 1.5% NaOH solution Although NaCN is not a precious commodity, the process can be cost-effective for concentrated electroplating baths. Removal of CN- was generally better by air stripping than by standard hypochlorite destruction. Addnl., 97.3-100% of the CN- was recovered by a NaOH bath, and the removal of Ce, Cu, Fe, Na, and In by S2- precipitation was equivalent to or better than the removal following hypochlorite destruction.

ANSWER 22 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1987:18.0609 CAPLUS

DOCUMENT NUMBER:

106:180609

TITLE:

Manufacture of low-copper steel from pig iron

containing copper

INVENTOR(S):

Tokuda, Masanori; Ito, Kimihisa; Obara, Yasushi;

Kawahara, Norizo

PATENT ASSIGNEE(S):

Iwate Steel Co., Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 7 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 61266515	Α	19861126	JP 1985-106915	19850521
PRIORITY APPLN. INFO.:			JP 1985-106915	19850521

AB Low-Cu steel is manufactured from molten pig iron containing Cu, C, and metal impurities by adding Na2S and Na2SO4 to transfer the Cu and impurities into slag. The vaporizing of Na is suppressed during the addition, but vaporized Na is recovered along with the Na2S and Na2SO4 in slag for reuse. A crucible was filled with 500 kg pig iron containing C 2.53, Si 0.80, Mn 0.53, P 0.115, S 0.135, Cu 1.43, Ti 0.105, Cr 0.062, V 0.040, and Zn 0.025% at $1400-1500^{\circ}$, 150 kg of a Na2S-Na2SO4 mixture (1:2) was added, and then slagged 3-5 min later. The adding-slagging process repeated 7 times and powdered CaC2 was then injected for desulfurization.

Molten steel containing C 0.32, Cu 0.03, and Si, Mn, P, Ti, Cr, V, and Zr <0.01% each was manufactured Vaporized Na2S (.apprx.15%) was recovered by scrubbing while in-slag Na2S and Na2SO4 were recovered by an eluting-precipitating

process.

L2 ANSWER 23 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1982:514848 CAPLUS

DOCUMENT NUMBER: 97:114848

TITLE: Treatment of mine wastewater PATENT ASSIGNEE(S): Toho Zinc Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	-			
JP 57081880	Α	19820522	JP 1980-157984	19801110
PRIORITY APPLN. INFO.:			JP 1980-157984	19801110

AB Mine drainage containing Fe, Zn, or other heavy metal ions is treated with a carbonized sawdust to remove Fe ion; then the effluent is treated with a sulfidizing agent and filtered through the carbonized sawdust to remove Zn ion. Thus, mine drainage containing 14.22 ppm Zn2+ and 0.042 ppm Cd2+ was passed through 6 carbonized sawdust columns in series, and a Na2S solution was fed to the 2 column. The removal was 100% for Cd and 97% for Zn.

L2 ANSWER 24 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1981:553011 CAPLUS

DOCUMENT NUMBER: 95:153011

TITLE: Zinc removal from aluminate solutions

INVENTOR(S): Bird, Robert D.; Vance, Harry R.

PATENT ASSIGNEE(S): Reynolds Metals Co., USA

SOURCE: U.S., 6 pp.

CODEN: USXXAM

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 4282191 A 19810804 US 1980-172420 19800725

PRIORITY APPLN. INFO.: US 1980-172420 A 19800725

AB The amount of Zn impurity in a caustic Na aluminate solution for preparing Al203

is decreased by adding ZnS seed crystals in the presence of S2-. The ZnS seed crystals are prepared by treating a caustic Na2ZnO2 solution with a sufficient amount of Na2S to form ZnS and have an excess of Na2S in an amount sufficient to give ≥0.150 g Na2S/L when the solution is added to the process liquor. The preceding solution is added to the Zn-containing process liquor to precipitate ZnS and give a solution after filtration containing <10 mg ZnO/L.

L2 ANSWER 25 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1978:448469 CAPLUS

DOCUMENT NUMBER:

89:48469

TITLE:

Heavy metal removal from wastewaters

INVENTOR(S):

Hirose, Kiyoshi; Fukatsu, Ryoki; Fujita, Takeshi Sakai Chemical Industry Co., Ltd., Sakai, Japan; S. K.

PATENT ASSIGNEE(S): Sakai Chemical Indus Engineering K. K.

SOURCE:

Jpn. Kokai Tokkyo Koho, 11 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 52126058 RITY APPLN. INFO.:	Α	19771022	JP 1975-48044 JP 1975-48044 A	19750418 19750418

AB Wastewaters are treated alkaline, then the heavy metals are precipitated as sulfides,

and the excess S2- in the wastewaters is precipitated. Thus, a wastewater containing

100 ppm each of Fe2+, Pb2+, and Zn2+ was heated at 70° , then NaOH (to pH 9) and 1% Na2S (1.5 equiv of the metals) was added to contain 0.9, 0.5, and <0.1 ppm, resp. in the supernatant. ZnCl2 (0.8 equivalent) was added to remove excess S2-.

L2 ANSWER 26 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1977:18485 CAPLUS

DOCUMENT NUMBER:

86:18485

TITLE:

Cadmium pigments with low acid-soluble cadmium content

INVENTOR(S):

Berger, Rolf; Kuch, Wolfgang; Endriss, Hartmut BASF Farben und Fasern A.-G., Fed. Rep. Ger.

PATENT ASSIGNEE(S):

Belg., 9 pp.

SOURCE: Belg., 9 p

DOCUMENT TYPE:

CODEN: BEXXAL

DOCUMENT II

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
				-	
BE 834804	A1	19760216	BE 1975-161191		19751023
DE 2513964	A1	19760930	DE 1975-2513964	,	19750329
FR 2293477	A1	19760702	FR 1975-32292		19751022
JP 51068630	Α	19760614	JP 1975-126951		19751023
NL 7513013	Α	19760609	NL 1975-13013		19751106
US 3977895	Α	19760831	US 1975-635720		19751126
BR 7508012	Α	19760824	BR 1975-8012		19751203
PRIORITY APPLN. INFO.:			DE 1974-2457390	Α	19741205
	,		DE 1975-2513964	Α	19750329

AB The content of acid-soluble Cd salts in red and yellow pigments was reduced by pretreating the pigments with SnCl4, then either adding Na2S to precipitate the Sn, and washing the pigment free of Cl, or neutralizing the pretreated mixture with NaOH, and adding ZnSO4 in HOAc, followed by aqueous Na2S. Typically, Cd red pigment in water was treated with aqueous SnCl4, followed by aqueous Na2S, washed, filtered, dried, and milled to give a pigment containing 0.0008% acid-soluble Cd salts rather than the 0.06% content of untreated

pigment.

ANSWER 27 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1976:109050 CAPLUS

DOCUMENT NUMBER:

84:109050

TITLE:

Removal of cobalt from zinc solutions

INVENTOR(S):

Neverov, L. P.; Chernova, R. A.

PATENT ASSIGNEE(S):

Ust-Kamenogorsk Lead-Zinc Combine, USSR

SOURCE:

U.S.S.R. From: Otkrytiya, Izobret., Prom. Obraztsy,

Tovarnye Znaki 1975, 52(40), 67.

CODEN: URXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Russian

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
SU 489798	A1	19751030	SU 1974-2040533	19740705
PRIORITY APPLN. INFO.:			SU 1974-2040533 A	19740705
AB Alkali metal xanthat	te was	used as the	precipitating reagent f	or the
purification of Zn				

solns. To decrease the consumption of the reagent and to facilitate the filtration of purified solns., alkali metal xanthate was mixed with alkali metal sulfide (Na2S) 1 weight part per 0.5-20 weight parts of xanthate.

ANSWER 28 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1969:515415 CAPLUS

DOCUMENT NUMBER:

71:115415

ORIGINAL REFERENCE NO.:

71:21509a,21512a

TITLE:

Purifying Bayer digester slurry

INVENTOR(S):

Hrishikesan, Kizhakke G.; Kane, James F.; Teas, Edward

B.; Brown, William Haddon

PATENT ASSIGNEE(S):

Reynolds Metals Co. U.S., 3 pp.

SOURCE:

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 3469935	. A	19690930	US 1965-456817	19650518
PRIORITY APPLN. INFO.:			US 1965-456817 A	19650518

AB To meet the purity specifications for metallic Al which limit the Zn content to $\leq 0.03\%$, the Bayer-process liquor should contain ≤0.012 g. ZnO/kg. In using Caribbean bauxite ores which are relatively high in Zn, the Bayer liquor may contain as much as 0.024 g. ZnO/kg. In the present process, an alkali metal sulfide is added to the digester slurry to precipitate ZnS, and the mixture is held in a quiescent

for 15-45 min. The resulting clarified liquor can be purified further by filtering through a bed of red mud. Thus, to digester slurry containing 93 q. Al203/kg. and 0.024 g. ZnO/kg. was added 0.18 g. Na25/kg. The mixture was held quiescent at 200°F. for 30 min. The supernatant liquor contained 0.01 g. ZnO/kg. Filtration through red mud further decreased

the ZnO to 0.006 g./kg. The Na2S for this process can be prepared by carbothermic reduction of a mixture of Na2SO4, Na2CO3, and NaAlO2 obtained as a by-product in the Bayer process.

ANSWER 29 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN L2

ACCESSION NUMBER:

1969:504154 CAPLUS

DOCUMENT NUMBER:

71:104154

ORIGINAL REFERENCE NO.: 71:19439a,19442a

TITLE:

Removal of zinc from Bayer process

liquors with sodium sulfide

AUTHOR(S):

Hansen, Robert Marius

CORPORATE SOURCE:

Kaiser Alum. and Chem. Corp., Gramercy, LA, USA

SOURCE:

Journal of Metals (1969), 21(9), 32-4

CODEN: JOMTAA; ISSN: 0148-6608

DOCUMENT TYPE:

Journal · English

LANGUAGE:

AB Some of the Zn present in Jamaican bauxite dissolves in the Bayer process and contaminates the finished Al2O3. To correct this, NaHS (a by-product of petroleum refining) is added to the hot Na aluminate liquor after sedimentation, but prior to clear-pressing. The Zn is precipitated as ZnS and removed at the clear press. The S is oxidized to Na2SO4 and accumulates in the liquor. However, the SO4 - enters the desilication product on subsequent cycles, so that the liquors need not be purged of Na2SO4.

anal. control method was developed, based on the principle of atomic absorption. Graphs show the solubility of Zn vs. Na2S at various initial Zn levels, the solubility of Zn vs. contact time at various Na2S levels, and Na2SO4 substitution in desilication product vs. Na2SO4 concentration

There is some insol. Zn in the ore, and some Na2S is recycled; hence an

ANSWER 30 OF 30 CAPLUS COPYRIGHT 2007 ACS on STN L2

ACCESSION NUMBER:

1969:432594 CAPLUS

DOCUMENT NUMBER:

71:32594

ORIGINAL REFERENCE NO.:

71:6039a

TITLE:

Purifying aluminate solutions

INVENTOR(S):

Hrishikesan, Kizhakke G.

PATENT ASSIGNEE(S):

Reynolds Metals Co.

SOURCE:

U.S., 3 pp.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 3445186	Α	19690520	US 1966-600377	19661209
PRIORITY APPLN. INFO.:			US 1966-600377 A	19661209

AB Dissolved Zn is removed from Bayer-process liquor by adding Na2S, whereby In is precipitated as InS. Thus, Bayer digestion slurry from the treatment of Haiti bauxite (Al203 45.86, Fe203 20.96, and ZnO 0.031%) contained Al203 93 and ZnO 0.024 g./kg. To this slurry at $200^{\circ}F$. was added 0.18 g. Na2S/kg., the mixture was held 60 min., starch was added to aid sedimentation, and the solids were allowed to settle. The clarified liquor contained 0.01 g. ZnO/kg.

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
•	ENTRY	SESSION
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FULL ESTIMATED COST	0.30	101.36
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	0.00	-23.40

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